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(54) **Title:** COMPOSITIONS AND METHODS FOR THERAPY AND DIAGNOSIS OF COLON CANCER

(57) **Abstract:** Compositions and methods for the therapy and diagnosis of cancer, such as colon cancer, are disclosed. Compositions may comprise one or more colon tumor proteins, immunogenic portions thereof, or polynucleotides that encode such portions. Alternatively, a therapeutic composition may comprise an antigen presenting cell that expresses a colon tumor protein, or a T cell that is specific for cells expressing such a protein. Such compositions may be used, for example, for the prevention and treatment of diseases such as colon cancer. Diagnostic methods based on detecting a colon tumor protein, or mRNA encoding such a protein, in a sample are also provided.

COMPOSITIONS AND METHODS FOR THERAPY AND DIAGNOSIS OF COLON CANCER

TECHNICAL FIELD

The present invention relates generally to therapy and diagnosis of
5 cancer, such as colon cancer. The invention is more specifically related to polypeptides
comprising at least a portion of a colon tumor protein, and to polynucleotides encoding
such polypeptides. Such polypeptides and polynucleotides may be used in vaccines and
pharmaceutical compositions for prevention and treatment of colon cancers, and for the
diagnosis and monitoring of such cancers.

10 BACKGROUND OF THE INVENTION

Cancer is a significant health problem throughout the world. Although
advances have been made in detection and therapy of cancer, no vaccine or other
universally successful method for prevention or treatment is currently available.
Current therapies, which are generally based on a combination of chemotherapy or
15 surgery and radiation, continue to prove inadequate in many patients.

Colon cancer is the second most frequently diagnosed malignancy in the
United States as well as the second most common cause of cancer death. An estimated
95,600 new cases of colon cancer will be diagnosed in 1998, with an estimated 47,700
deaths. The five-year survival rate for patients with colorectal cancer detected in an
20 early localized stage is 92%; unfortunately, only 37% of colorectal cancer is diagnosed
at this stage. The survival rate drops to 64% if the cancer is allowed to spread to
adjacent organs or lymph nodes, and to 7% in patients with distant metastases.

The prognosis of colon cancer is directly related to the degree of
penetration of the tumor through the bowel wall and the presence or absence of nodal
25 involvement, consequently, early detection and treatment are especially important.
Currently, diagnosis is aided by the use of screening assays for fecal occult blood,
sigmoidoscopy, colonoscopy and double contrast barium enemas. Treatment regimens
are determined by the type and stage of the cancer, and include surgery, radiation
therapy and/or chemotherapy. Recurrence following surgery (the most common form

of therapy) is a major problem and is often the ultimate cause of death. In spite of considerable research into therapies for the disease, colon cancer remains difficult to diagnose and treat. In spite of considerable research into therapies for these and other cancers, colon cancer remains difficult to diagnose and treat effectively. Accordingly, there is a need in the art for improved methods for detecting and treating such cancers. The present invention fulfills these needs and further provides other related advantages.

SUMMARY OF THE INVENTION

Briefly stated, the present invention provides compositions and methods for the diagnosis and therapy of cancer, such as colon cancer. In one aspect, the present invention provides polypeptides comprising at least a portion of a colon tumor protein, or a variant thereof. Certain portions and other variants are immunogenic, such that the ability of the variant to react with antigen-specific antisera is not substantially diminished. Within certain embodiments, the polypeptide comprises a sequence that is encoded by a polynucleotide sequence selected from the group consisting of: (a) sequences recited in SEQ ID NOs:1-1556; (b) variants of a sequence recited in SEQ ID NO: 1-1556; and (c) complements of a sequence of (a) or (b).

The present invention further provides polynucleotides that encode a polypeptide as described above, or a portion thereof (such as a portion encoding at least 15 amino acid residues of a colon tumor protein), expression vectors comprising such polynucleotides and host cells transformed or transfected with such expression vectors.

Within other aspects, the present invention provides pharmaceutical compositions comprising a polypeptide or polynucleotide as described above and a physiologically acceptable carrier.

Within a related aspect of the present invention, vaccines for prophylactic or therapeutic use are provided. Such vaccines comprise a polypeptide or polynucleotide as described above and an immunostimulant.

The present invention further provides pharmaceutical compositions that comprise: (a) an antibody or antigen-binding fragment thereof that specifically binds to a colon tumor protein; and (b) a physiologically acceptable carrier.

Within further aspects, the present invention provides pharmaceutical compositions comprising: (a) an antigen presenting cell that expresses a polypeptide as described above and (b) a pharmaceutically acceptable carrier or excipient. Antigen presenting cells include dendritic cells, macrophages, monocytes, fibroblasts and B cells.

Within related aspects, vaccines are provided that comprise: (a) an antigen presenting cell that expresses a polypeptide as described above and (b) an immunostimulant.

The present invention further provides, in other aspects, fusion proteins that comprise at least one polypeptide as described above, as well as polynucleotides encoding such fusion proteins.

Within related aspects, pharmaceutical compositions comprising a fusion protein, or a polynucleotide encoding a fusion protein, in combination with a physiologically acceptable carrier are provided.

Vaccines are further provided, within other aspects, that comprise a fusion protein, or a polynucleotide encoding a fusion protein, in combination with an immunostimulant.

Within further aspects, the present invention provides methods for inhibiting the development of a cancer in a patient, comprising administering to a patient a pharmaceutical composition or vaccine as recited above. The patient may be afflicted with colon cancer, in which case the methods provide treatment for the disease, or patient considered at risk for such a disease may be treated prophylactically.

The present invention further provides, within other aspects, methods for removing tumor cells from a biological sample, comprising contacting a biological sample with T cells that specifically react with a colon tumor protein, wherein the step of contacting is performed under conditions and for a time sufficient to permit the removal of cells expressing the protein from the sample.

Within related aspects, methods are provided for inhibiting the development of a cancer in a patient, comprising administering to a patient a biological sample treated as described above.

Methods are further provided, within other aspects, for stimulating and/or expanding T cells specific for a colon tumor protein, comprising contacting T cells with one or more of: (i) a polypeptide as described above; (ii) a polynucleotide encoding such a polypeptide; and/or (iii) an antigen presenting cell that expresses such a polypeptide; under conditions and for a time sufficient to permit the stimulation and/or expansion of T cells. Isolated T cell populations comprising T cells prepared as described above are also provided.

Within further aspects, the present invention provides methods for inhibiting the development of a cancer in a patient, comprising administering to a patient an effective amount of a T cell population as described above.

The present invention further provides methods for inhibiting the development of a cancer in a patient, comprising the steps of: (a) incubating CD4⁺ and/or CD8⁺ T cells isolated from a patient with one or more of: (i) a polypeptide comprising at least an immunogenic portion of a colon tumor protein; (ii) a polynucleotide encoding such a polypeptide; and (iii) an antigen-presenting cell that expressed such a polypeptide; and (b) administering to the patient an effective amount of the proliferated T cells, and thereby inhibiting the development of a cancer in the patient. Proliferated cells may, but need not, be cloned prior to administration to the patient.

Within further aspects, the present invention provides methods for determining the presence or absence of a cancer in a patient, comprising: (a) contacting a biological sample obtained from a patient with a binding agent that binds to a polypeptide as recited above; (b) detecting in the sample an amount of polypeptide that binds to the binding agent; and (c) comparing the amount of polypeptide with a predetermined cut-off value, and therefrom determining the presence or absence of a cancer in the patient. Within preferred embodiments, the binding agent is an antibody, more preferably a monoclonal antibody. The cancer may be colon cancer.

The present invention also provides, within other aspects, methods for monitoring the progression of a cancer in a patient. Such methods comprise the steps of: (a) contacting a biological sample obtained from a patient at a first point in time with a binding agent that binds to a polypeptide as recited above; (b) detecting in the

- sample an amount of polypeptide that binds to the binding agent; (c) repeating steps (a) and (b) using a biological sample obtained from the patient at a subsequent point in time; and (d) comparing the amount of polypeptide detected in step (c) with the amount detected in step (b) and therefrom monitoring the progression of the cancer in the patient.

The present invention further provides, within other aspects, methods for determining the presence or absence of a cancer in a patient, comprising the steps of: (a) contacting a biological sample obtained from a patient with an oligonucleotide that hybridizes to a polynucleotide that encodes a colon tumor protein; (b) detecting in the sample a level of a polynucleotide, preferably mRNA, that hybridizes to the oligonucleotide; and (c) comparing the level of polynucleotide that hybridizes to the oligonucleotide with a predetermined cut-off value, and therefrom determining the presence or absence of a cancer in the patient. Within certain embodiments, the amount of mRNA is detected via polymerase chain reaction using, for example, at least one oligonucleotide primer that hybridizes to a polynucleotide encoding a polypeptide as recited above, or a complement of such a polynucleotide. Within other embodiments, the amount of mRNA is detected using a hybridization technique, employing an oligonucleotide probe that hybridizes to a polynucleotide that encodes a polypeptide as recited above, or a complement of such a polynucleotide.

- In related aspects, methods are provided for monitoring the progression of a cancer in a patient, comprising the steps of: (a) contacting a biological sample obtained from a patient with an oligonucleotide that hybridizes to a polynucleotide that encodes a colon tumor protein; (b) detecting in the sample an amount of a polynucleotide that hybridizes to the oligonucleotide; (c) repeating steps (a) and (b) using a biological sample obtained from the patient at a subsequent point in time; and (d) comparing the amount of polynucleotide detected in step (c) with the amount detected in step (b) and therefrom monitoring the progression of the cancer in the patient.

- Within further aspects, the present invention provides antibodies, such as monoclonal antibodies, that bind to a polypeptide as described above, as well as

diagnostic kits comprising such antibodies. Diagnostic kits comprising one or more oligonucleotide probes or primers as described above are also provided.

These and other aspects of the present invention will become apparent upon reference to the following detailed description. All references disclosed herein are hereby incorporated by reference in their entirety as if each was incorporated individually.

DETAILED DESCRIPTION OF THE INVENTION

As noted above, the present invention is generally directed to compositions and methods for the therapy and diagnosis of cancer, such as colon cancer. The compositions described herein may include colon tumor polypeptides, polynucleotides encoding such polypeptides, binding agents such as antibodies, antigen presenting cells (APCs) and/or immune system cells (*e.g.*, T cells). Polypeptides of the present invention generally comprise at least a portion (such as an immunogenic portion) of a colon tumor protein or a variant thereof. A "colon tumor protein" is a protein that is expressed in colon tumor cells at a level that is at least two fold, and preferably at least five fold, greater than the level of expression in a normal tissue, as determined using a representative assay provided herein. Certain colon tumor proteins are tumor proteins that react detectably (within an immunoassay, such as an ELISA or Western blot) with antisera of a patient afflicted with colon cancer. Polynucleotides of the subject invention generally comprise a DNA or RNA sequence that encodes all or a portion of such a polypeptide, or that is complementary to such a sequence. Antibodies are generally immune system proteins, or antigen-binding fragments thereof, that are capable of binding to a polypeptide as described above. Antigen presenting cells include dendritic cells, macrophages, monocytes, fibroblasts and B-cells that express a polypeptide as described above. T cells that may be employed within such compositions are generally T cells that are specific for a polypeptide as described above.

The present invention is based on the discovery of human colon tumor proteins. Sequences of polynucleotides encoding specific tumor proteins are provided in SEQ ID NOs:1-1556.

COLON TUMOR PROTEIN POLYNUCLEOTIDES

Any polynucleotide that encodes a colon tumor protein or a portion or other variant thereof as described herein is encompassed by the present invention. Preferred polynucleotides comprise at least 15 consecutive nucleotides, preferably at least 30 consecutive nucleotides and more preferably at least 45 consecutive nucleotides, that encode a portion of a colon tumor protein. More preferably, a polynucleotide encodes an immunogenic portion of a colon tumor protein. Polynucleotides complementary to any such sequences are also encompassed by the present invention. Polynucleotides may be single-stranded (coding or antisense) or double-stranded, and may be DNA (genomic, cDNA or synthetic) or RNA molecules. RNA molecules include HnRNA molecules, which contain introns and correspond to a DNA molecule in a one-to-one manner, and mRNA molecules, which do not contain introns. Additional coding or non-coding sequences may, but need not, be present within a polynucleotide of the present invention, and a polynucleotide may, but need not, be linked to other molecules and/or support materials.

Polynucleotides may comprise a native sequence (*i.e.*, an endogenous sequence that encodes a colon tumor protein or a portion thereof) or may comprise a variant of such a sequence. Polynucleotide variants may contain one or more substitutions, additions, deletions and/or insertions such that the immunogenicity of the encoded polypeptide is not diminished, relative to a native tumor protein. The effect on the immunogenicity of the encoded polypeptide may generally be assessed as described herein. Variants preferably exhibit at least about 70% identity, more preferably at least about 80% identity and most preferably at least about 90% identity to a polynucleotide sequence that encodes a native colon tumor protein or a portion thereof. The term “variants” also encompasses homologous genes of xenogenic origin.

Two polynucleotide or polypeptide sequences are said to be “identical” if the sequence of nucleotides or amino acids in the two sequences is the same when aligned for maximum correspondence as described below. Comparisons between two sequences are typically performed by comparing the sequences over a comparison window to identify and compare local regions of sequence similarity. A “comparison window” as used herein, refers to a segment of at least about 20 contiguous positions,

usually 30 to about 75, 40 to about 50, in which a sequence may be compared to a reference sequence of the same number of contiguous positions after the two sequences are optimally aligned.

- Optimal alignment of sequences for comparison may be conducted using
- 5 the Megalign program in the Lasergene suite of bioinformatics software (DNASTAR, Inc., Madison, WI), using default parameters. This program embodies several alignment schemes described in the following references: Dayhoff, M.O. (1978) A model of evolutionary change in proteins – Matrices for detecting distant relationships. In Dayhoff, M.O. (ed.) *Atlas of Protein Sequence and Structure*, National Biomedical
 - 10 Research Foundation, Washington DC Vol. 5, Suppl. 3, pp. 345-358; Hein J. (1990) Unified Approach to Alignment and Phylogenies pp. 626-645 *Methods in Enzymology* vol. 183, Academic Press, Inc., San Diego, CA; Higgins, D.G. and Sharp, P.M. (1989) *CABIOS* 5:151-153; Myers, E.W. and Muller W. (1988) *CABIOS* 4:11-17; Robinson, E.D. (1971) *Comb. Theor* 11:105; Santou, N. Nes, M. (1987) *Mol. Biol. Evol.* 4:406-
 - 15 425; Sneath, P.H.A. and Sokal, R.R. (1973) *Numerical Taxonomy – the Principles and Practice of Numerical Taxonomy*, Freeman Press, San Francisco, CA; Wilbur, W.J. and Lipman, D.J. (1983) *Proc. Natl. Acad., Sci. USA* 80:726-730.

Preferably, the “percentage of sequence identity” is determined by comparing two optimally aligned sequences over a window of comparison of at least 20

- 20 positions, wherein the portion of the polynucleotide or polypeptide sequence in the comparison window may comprise additions or deletions (*i.e.*, gaps) of 20 percent or less, usually 5 to 15 percent, or 10 to 12 percent, as compared to the reference sequences (which does not comprise additions or deletions) for optimal alignment of the two sequences. The percentage is calculated by determining the number of positions at
- 25 which the identical nucleic acid bases or amino acid residue occurs in both sequences to yield the number of matched positions, dividing the number of matched positions by the total number of positions in the reference sequence (*i.e.*, the window size) and multiplying the results by 100 to yield the percentage of sequence identity.

Variants may also, or alternatively, be substantially homologous to a

- 30 native gene, or a portion or complement thereof. Such polynucleotide variants are capable of hybridizing under moderately stringent conditions to a naturally occurring

DNA sequence encoding a native colon tumor protein (or a complementary sequence). Suitable moderately stringent conditions include prewashing in a solution of 5 X SSC, 0.5% SDS, 1.0 mM EDTA (pH 8.0); hybridizing at 50°C-65°C, 5 X SSC, overnight; followed by washing twice at 65°C for 20 minutes with each of 2X, 0.5X and 0.2X SSC
5 containing 0.1% SDS.

It will be appreciated by those of ordinary skill in the art that, as a result of the degeneracy of the genetic code, there are many nucleotide sequences that encode a polypeptide as described herein. Some of these polynucleotides bear minimal homology to the nucleotide sequence of any native gene. Nonetheless, polynucleotides
10 that vary due to differences in codon usage are specifically contemplated by the present invention. Further, alleles of the genes comprising the polynucleotide sequences provided herein are within the scope of the present invention. Alleles are endogenous genes that are altered as a result of one or more mutations, such as deletions, additions and/or substitutions of nucleotides. The resulting mRNA and protein may, but need
15 not, have an altered structure or function. Alleles may be identified using standard techniques (such as hybridization, amplification and/or database sequence comparison).

Polynucleotides may be prepared using any of a variety of techniques. For example, a polynucleotide may be identified, as described in more detail below, by screening a microarray of cDNAs for tumor-associated expression (*i.e.*, expression that
20 is at least two fold greater in a colon tumor than in normal tissue, as determined using a representative assay provided herein). Such screens may be performed using a Synteni microarray (Palo Alto, CA) according to the manufacturer's instructions (and essentially as described by Schena et al., *Proc. Natl. Acad. Sci. USA* 93:10614-10619, 1996 and Heller et al., *Proc. Natl. Acad. Sci. USA* 94:2150-2155, 1997). Alternatively,
25 polynucleotides may be amplified from cDNA prepared from cells expressing the proteins described herein, such as colon tumor cells. Such polynucleotides may be amplified via polymerase chain reaction (PCR). For this approach, sequence-specific primers may be designed based on the sequences provided herein, and may be purchased or synthesized.

30 An amplified portion may be used to isolate a full length gene from a suitable library (*e.g.*, a colon tumor cDNA library) using well known techniques.

Within such techniques, a library (cDNA or genomic) is screened using one or more polynucleotide probes or primers suitable for amplification. Preferably, a library is size-selected to include larger molecules. Random primed libraries may also be preferred for identifying 5' and upstream regions of genes. Genomic libraries are preferred for obtaining introns and extending 5' sequences.

For hybridization techniques, a partial sequence may be labeled (*e.g.*, by nick-translation or end-labeling with ^{32}P) using well known techniques. A bacterial or bacteriophage library is then screened by hybridizing filters containing denatured bacterial colonies (or lawns containing phage plaques) with the labeled probe (*see* Sambrook et al., *Molecular Cloning: A Laboratory Manual*, Cold Spring Harbor Laboratories, Cold Spring Harbor, NY, 1989). Hybridizing colonies or plaques are selected and expanded, and the DNA is isolated for further analysis. cDNA clones may be analyzed to determine the amount of additional sequence by, for example, PCR using a primer from the partial sequence and a primer from the vector. Restriction maps and partial sequences may be generated to identify one or more overlapping clones. The complete sequence may then be determined using standard techniques, which may involve generating a series of deletion clones. The resulting overlapping sequences are then assembled into a single contiguous sequence. A full length cDNA molecule can be generated by ligating suitable fragments, using well known techniques.

Alternatively, there are numerous amplification techniques for obtaining a full length coding sequence from a partial cDNA sequence. Within such techniques, amplification is generally performed via PCR. Any of a variety of commercially available kits may be used to perform the amplification step. Primers may be designed using, for example, software well known in the art. Primers are preferably 22-30 nucleotides in length, have a GC content of at least 50% and anneal to the target sequence at temperatures of about 68°C to 72°C. The amplified region may be sequenced as described above, and overlapping sequences assembled into a contiguous sequence.

One such amplification technique is inverse PCR (*see* Triglia et al., *Nucl. Acids Res.* 16:8186, 1988), which uses restriction enzymes to generate a fragment in the known region of the gene. The fragment is then circularized by intramolecular ligation

and used as a template for PCR with divergent primers derived from the known region. Within an alternative approach, sequences adjacent to a partial sequence may be retrieved by amplification with a primer to a linker sequence and a primer specific to a known region. The amplified sequences are typically subjected to a second round of
5 amplification with the same linker primer and a second primer specific to the known region. A variation on this procedure, which employs two primers that initiate extension in opposite directions from the known sequence, is described in WO 96/38591. Another such technique is known as "rapid amplification of cDNA ends" or RACE. This technique involves the use of an internal primer and an external primer,
10 which hybridizes to a polyA region or vector sequence, to identify sequences that are 5' and 3' of a known sequence. Additional techniques include capture PCR (Lagerstrom et al., *PCR Methods Applic. 1*:111-19, 1991) and walking PCR (Parker et al., *Nucl. Acids. Res. 19*:3055-60, 1991). Other methods employing amplification may also be employed to obtain a full length cDNA sequence.

15 In certain instances, it is possible to obtain a full length cDNA sequence by analysis of sequences provided in an expressed sequence tag (EST) database, such as that available from GenBank. Searches for overlapping ESTs may generally be performed using well known programs (e.g., NCBI BLAST searches), and such ESTs may be used to generate a contiguous full length sequence. Full length DNA sequences
20 may also be obtained by analysis of genomic fragments.

Certain nucleic acid sequences of cDNA molecules encoding portions of colon tumor proteins are provided in SEQ ID NOs: 1-1556.

Polynucleotide variants may generally be prepared by any method known in the art, including chemical synthesis by, for example, solid phase
25 phosphoramidite chemical synthesis. Modifications in a polynucleotide sequence may also be introduced using standard mutagenesis techniques, such as oligonucleotide-directed site-specific mutagenesis (see Adelman et al., *DNA 2*:183, 1983). Alternatively, RNA molecules may be generated by *in vitro* or *in vivo* transcription of DNA sequences encoding a colon tumor protein, or portion thereof, provided that the
30 DNA is incorporated into a vector with a suitable RNA polymerase promoter (such as T7 or SP6). Certain portions may be used to prepare an encoded polypeptide, as

described herein. In addition, or alternatively, a portion may be administered to a patient such that the encoded polypeptide is generated *in vivo* (e.g., by transfecting antigen-presenting cells, such as dendritic cells, with a cDNA construct encoding a colon tumor polypeptide, and administering the transfected cells to the patient).

- 5 A portion of a sequence complementary to a coding sequence (*i.e.*, an antisense polynucleotide) may also be used as a probe or to modulate gene expression. cDNA constructs that can be transcribed into antisense RNA may also be introduced into cells or tissues to facilitate the production of antisense RNA. An antisense polynucleotide may be used, as described herein, to inhibit expression of a tumor
- 10 protein. Antisense technology can be used to control gene expression through triple-helix formation, which compromises the ability of the double helix to open sufficiently for the binding of polymerases, transcription factors or regulatory molecules (*see* Gee et al., *In Huber and Carr, Molecular and Immunologic Approaches*, Futura Publishing Co. (Mt. Kisco, NY; 1994)). Alternatively, an antisense molecule may be designed to
- 15 hybridize with a control region of a gene (e.g., promoter, enhancer or transcription initiation site), and block transcription of the gene; or to block translation by inhibiting binding of a transcript to ribosomes.

- A portion of a coding sequence, or of a complementary sequence, may also be designed as a probe or primer to detect gene expression. Probes may be labeled
- 20 with a variety of reporter groups, such as radionuclides and enzymes, and are preferably at least 10 nucleotides in length, more preferably at least 20 nucleotides in length and still more preferably at least 30 nucleotides in length. Primers, as noted above, are preferably 22-30 nucleotides in length.

- Any polynucleotide may be further modified to increase stability *in vivo*.
- 25 Possible modifications include, but are not limited to, the addition of flanking sequences at the 5' and/or 3' ends; the use of phosphorothioate or 2' O-methyl rather than phosphodiesterase linkages in the backbone; and/or the inclusion of nontraditional bases such as inosine, queosine and wybutosine, as well as acetyl- methyl-, thio- and other modified forms of adenine, cytidine, guanine, thymine and uridine.

- 30 Nucleotide sequences as described herein may be joined to a variety of other nucleotide sequences using established recombinant DNA techniques. For

example, a polynucleotide may be cloned into any of a variety of cloning vectors, including plasmids, phagemids, lambda phage derivatives and cosmids. Vectors of particular interest include expression vectors, replication vectors, probe generation vectors and sequencing vectors. In general, a vector will contain an origin of replication functional in at least one organism, convenient restriction endonuclease sites and one or more selectable markers. Other elements will depend upon the desired use, and will be apparent to those of ordinary skill in the art.

Within certain embodiments, polynucleotides may be formulated so as to permit entry into a cell of a mammal, and expression therein. Such formulations are particularly useful for therapeutic purposes, as described below. Those of ordinary skill in the art will appreciate that there are many ways to achieve expression of a polynucleotide in a target cell, and any suitable method may be employed. For example, a polynucleotide may be incorporated into a viral vector such as, but not limited to, adenovirus, adeno-associated virus, retrovirus, or vaccinia or other pox virus (*e.g.*, avian pox virus). The polynucleotides may also be administered as naked plasmid vectors. Techniques for incorporating DNA into such vectors are well known to those of ordinary skill in the art. A retroviral vector may additionally transfer or incorporate a gene for a selectable marker (to aid in the identification or selection of transduced cells) and/or a targeting moiety, such as a gene that encodes a ligand for a receptor on a specific target cell, to render the vector target specific. Targeting may also be accomplished using an antibody, by methods known to those of ordinary skill in the art.

Other formulations for therapeutic purposes include colloidal dispersion systems, such as macromolecule complexes, nanocapsules, microspheres, beads, and lipid-based systems including oil-in-water emulsions, micelles, mixed micelles, and liposomes. A preferred colloidal system for use as a delivery vehicle *in vitro* and *in vivo* is a liposome (*i.e.*, an artificial membrane vesicle). The preparation and use of such systems is well known in the art.

COLON TUMOR POLYPEPTIDES

Within the context of the present invention, polypeptides may comprise at least an immunogenic portion of a colon tumor protein or a variant thereof, as

described herein. As noted above, a "colon tumor protein" is a protein that is expressed by colon tumor cells. Proteins that are colon tumor proteins also react detectably within an immunoassay (such as an ELISA) with antisera from a patient with colon cancer. Polypeptides as described herein may be of any length. Additional sequences derived from the native protein and/or heterologous sequences may be present, and such sequences may (but need not) possess further immunogenic or antigenic properties.

An "immunogenic portion," as used herein is a portion of a protein that is recognized (*i.e.*, specifically bound) by a B-cell and/or T-cell surface antigen receptor. Such immunogenic portions generally comprise at least 5 amino acid residues, more preferably at least 10, and still more preferably at least 20 amino acid residues of a colon tumor protein or a variant thereof. Certain preferred immunogenic portions include peptides in which an N-terminal leader sequence and/or transmembrane domain have been deleted. Other preferred immunogenic portions may contain a small N- and/or C-terminal deletion (*e.g.*, 1-30 amino acids, preferably 5-15 amino acids), relative to the mature protein.

Immunogenic portions may generally be identified using well known techniques, such as those summarized in Paul, *Fundamental Immunology*, 3rd ed., 243-247 (Raven Press, 1993) and references cited therein. Such techniques include screening polypeptides for the ability to react with antigen-specific antibodies, antisera and/or T-cell lines or clones. As used herein, antisera and antibodies are "antigen-specific" if they specifically bind to an antigen (*i.e.*, they react with the protein in an ELISA or other immunoassay, and do not react detectably with unrelated proteins). Such antisera and antibodies may be prepared as described herein, and using well known techniques. An immunogenic portion of a native colon tumor protein is a portion that reacts with such antisera and/or T-cells at a level that is not substantially less than the reactivity of the full length polypeptide (*e.g.*, in an ELISA and/or T-cell reactivity assay). Such immunogenic portions may react within such assays at a level that is similar to or greater than the reactivity of the full length polypeptide. Such screens may generally be performed using methods well known to those of ordinary skill in the art, such as those described in Harlow and Lane, *Antibodies: A Laboratory Manual*, Cold Spring Harbor Laboratory, 1988. For example, a polypeptide may be

immobilized on a solid support and contacted with patient sera to allow binding of antibodies within the sera to the immobilized polypeptide. Unbound sera may then be removed and bound antibodies detected using, for example, ^{125}I -labeled Protein A.

As noted above, a composition may comprise a variant of a native colon
5 tumor protein. A polypeptide "variant," as used herein, is a polypeptide that differs from a native colon tumor protein in one or more substitutions, deletions, additions and/or insertions, such that the immunogenicity of the polypeptide is not substantially diminished. In other words, the ability of a variant to react with antigen-specific antisera may be enhanced or unchanged, relative to the native protein, or may be
10 diminished by less than 50%, and preferably less than 20%, relative to the native protein. Such variants may generally be identified by modifying one of the above polypeptide sequences and evaluating the reactivity of the modified polypeptide with antigen-specific antibodies or antisera as described herein. Preferred variants include those in which one or more portions, such as an N-terminal leader sequence or
15 transmembrane domain, have been removed. Other preferred variants include variants in which a small portion (e.g., 1-30 amino acids, preferably 5-15 amino acids) has been removed from the N- and/or C-terminal of the mature protein.

Polypeptide variants preferably exhibit at least about 70%, more preferably at least about 90% and most preferably at least about 95% identity
20 (determined as described above) to the identified polypeptides.

Preferably, a variant contains conservative substitutions. A "conservative substitution" is one in which an amino acid is substituted for another amino acid that has similar properties, such that one skilled in the art of peptide chemistry would expect the secondary structure and hydrophobic nature of the
25 polypeptide to be substantially unchanged. Amino acid substitutions may generally be made on the basis of similarity in polarity, charge, solubility, hydrophobicity, hydrophilicity and/or the amphipathic nature of the residues. For example, negatively charged amino acids include aspartic acid and glutamic acid; positively charged amino acids include lysine and arginine; and amino acids with uncharged polar head groups
30 having similar hydrophilicity values include leucine, isoleucine and valine; glycine and alanine; asparagine and glutamine; and serine, threonine, phenylalanine and tyrosine.

Other groups of amino acids that may represent conservative changes include: (1) ala, pro, gly, glu, asp, gln, asn, ser, thr; (2) cys, ser, tyr, thr; (3) val, ile, leu, met, ala, phe; (4) lys, arg, his; and (5) phe, tyr, trp, his. A variant may also, or alternatively, contain nonconservative changes. In a preferred embodiment, variant polypeptides differ from a native sequence by substitution, deletion or addition of five amino acids or fewer. Variants may also (or alternatively) be modified by, for example, the deletion or addition of amino acids that have minimal influence on the immunogenicity, secondary structure and hydrophobic nature of the polypeptide.

As noted above, polypeptides may comprise a signal (or leader) sequence at the N-terminal end of the protein, which co-translationally or post-translationally directs transfer of the protein. The polypeptide may also be conjugated to a linker or other sequence for ease of synthesis, purification or identification of the polypeptide (e.g., poly-His), or to enhance binding of the polypeptide to a solid support. For example, a polypeptide may be conjugated to an immunoglobulin Fc region.

Polypeptides may be prepared using any of a variety of well known techniques. Recombinant polypeptides encoded by DNA sequences as described above may be readily prepared from the DNA sequences using any of a variety of expression vectors known to those of ordinary skill in the art. Expression may be achieved in any appropriate host cell that has been transformed or transfected with an expression vector containing a DNA molecule that encodes a recombinant polypeptide. Suitable host cells include prokaryotes, yeast, and higher eukaryotic cells, such as mammalian cells and plant cells. Preferably, the host cells employed are *E. coli*, yeast or a mammalian cell line such as COS or CHO. Supernatants from suitable host/vector systems which secrete recombinant protein or polypeptide into culture media may be first concentrated using a commercially available filter. Following concentration, the concentrate may be applied to a suitable purification matrix such as an affinity matrix or an ion exchange resin. Finally, one or more reverse phase HPLC steps can be employed to further purify a recombinant polypeptide.

Portions and other variants having less than about 100 amino acids, and generally less than about 50 amino acids, may also be generated by synthetic means, using techniques well known to those of ordinary skill in the art. For example, such

polypeptides may be synthesized using any of the commercially available solid-phase techniques, such as the Merrifield solid-phase synthesis method, where amino acids are sequentially added to a growing amino acid chain. *See* Merrifield, *J. Am. Chem. Soc.* 85:2149-2146, 1963. Equipment for automated synthesis of polypeptides is commercially available from suppliers such as Perkin Elmer/Applied BioSystems Division (Foster City, CA), and may be operated according to the manufacturer's instructions.

Within certain specific embodiments, a polypeptide may be a fusion protein that comprises multiple polypeptides as described herein, or that comprises at least one polypeptide as described herein and an unrelated sequence, such as a known tumor protein. A fusion partner may, for example, assist in providing T helper epitopes (an immunological fusion partner), preferably T helper epitopes recognized by humans, or may assist in expressing the protein (an expression enhancer) at higher yields than the native recombinant protein. Certain preferred fusion partners are both immunological and expression enhancing fusion partners. Other fusion partners may be selected so as to increase the solubility of the protein or to enable the protein to be targeted to desired intracellular compartments. Still further fusion partners include affinity tags, which facilitate purification of the protein.

Fusion proteins may generally be prepared using standard techniques, including chemical conjugation. Preferably, a fusion protein is expressed as a recombinant protein, allowing the production of increased levels, relative to a non-fused protein, in an expression system. Briefly, DNA sequences encoding the polypeptide components may be assembled separately, and ligated into an appropriate expression vector. The 3' end of the DNA sequence encoding one polypeptide component is ligated, with or without a peptide linker, to the 5' end of a DNA sequence encoding the second polypeptide component so that the reading frames of the sequences are in phase. This permits translation into a single fusion protein that retains the biological activity of both component polypeptides.

A peptide linker sequence may be employed to separate the first and second polypeptide components by a distance sufficient to ensure that each polypeptide folds into its secondary and tertiary structures. Such a peptide linker sequence is

incorporated into the fusion protein using standard techniques well known in the art. Suitable peptide linker sequences may be chosen based on the following factors: (1) their ability to adopt a flexible extended conformation; (2) their inability to adopt a secondary structure that could interact with functional epitopes on the first and second polypeptides; and (3) the lack of hydrophobic or charged residues that might react with the polypeptide functional epitopes. Preferred peptide linker sequences contain Gly, Asn and Ser residues. Other near neutral amino acids, such as Thr and Ala may also be used in the linker sequence. Amino acid sequences which may be usefully employed as linkers include those disclosed in Maratea et al., *Gene* 40:39-46, 1985; Murphy et al., *Proc. Natl. Acad. Sci. USA* 83:8258-8262, 1986; U.S. Patent No. 4,935,233 and U.S. Patent No. 4,751,180. The linker sequence may generally be from 1 to about 50 amino acids in length. Linker sequences are not required when the first and second polypeptides have non-essential N-terminal amino acid regions that can be used to separate the functional domains and prevent steric interference.

The ligated DNA sequences are operably linked to suitable transcriptional or translational regulatory elements. The regulatory elements responsible for expression of DNA are located only 5' to the DNA sequence encoding the first polypeptides. Similarly, stop codons required to end translation and transcription termination signals are only present 3' to the DNA sequence encoding the second polypeptide.

Fusion proteins are also provided. Such proteins comprise a polypeptide as described herein together with an unrelated immunogenic protein. Preferably the immunogenic protein is capable of eliciting a recall response. Examples of such proteins include tetanus, tuberculosis and hepatitis proteins (*see*, for example, Stoute et al. *New Engl. J. Med.*, 336:86-91, 1997).

Within preferred embodiments, an immunological fusion partner is derived from protein D, a surface protein of the gram-negative bacterium *Haemophilus influenza B* (WO 91/18926). Preferably, a protein D derivative comprises approximately the first third of the protein (*e.g.*, the first N-terminal 100-110 amino acids), and a protein D derivative may be lipidated. Within certain preferred embodiments, the first 109 residues of a Lipoprotein D fusion partner is included on the

N-terminus to provide the polypeptide with additional exogenous T-cell epitopes and to increase the expression level in *E. coli* (thus functioning as an expression enhancer). The lipid tail ensures optimal presentation of the antigen to antigen presenting cells. Other fusion partners include the non-structural protein from influenzae virus, NS1 (hemagglutinin). Typically, the N-terminal 81 amino acids are used, although different fragments that include T-helper epitopes may be used.

In another embodiment, the immunological fusion partner is the protein known as LYTA, or a portion thereof (preferably a C-terminal portion). LYTA is derived from *Streptococcus pneumoniae*, which synthesizes an N-acetyl-L-alanine amidase known as amidase LYTA (encoded by the *LytA* gene; *Gene* 43:265-292, 1986). LYTA is an autolysin that specifically degrades certain bonds in the peptidoglycan backbone. The C-terminal domain of the LYTA protein is responsible for the affinity to the choline or to some choline analogues such as DEAE. This property has been exploited for the development of *E. coli* C-LYTA expressing plasmids useful for expression of fusion proteins. Purification of hybrid proteins containing the C-LYTA fragment at the amino terminus has been described (see *Biotechnology* 10:795-798, 1992). Within a preferred embodiment, a repeat portion of LYTA may be incorporated into a fusion protein. A repeat portion is found in the C-terminal region starting at residue 178. A particularly preferred repeat portion incorporates residues 188-305.

In general, polypeptides (including fusion proteins) and polynucleotides as described herein are isolated. An "isolated" polypeptide or polynucleotide is one that is removed from its original environment. For example, a naturally-occurring protein is isolated if it is separated from some or all of the coexisting materials in the natural system. Preferably, such polypeptides are at least about 90% pure, more preferably at least about 95% pure and most preferably at least about 99% pure. A polynucleotide is considered to be isolated if, for example, it is cloned into a vector that is not a part of the natural environment.

BINDING AGENTS

The present invention further provides agents, such as antibodies and antigen-binding fragments thereof, that specifically bind to a colon tumor protein. As used herein, an antibody, or antigen-binding fragment thereof, is said to "specifically bind" to a colon tumor protein if it reacts at a detectable level (within, for example, an ELISA) with a colon tumor protein, and does not react detectably with unrelated proteins under similar conditions. As used herein, "binding" refers to a noncovalent association between two separate molecules such that a complex is formed. The ability to bind may be evaluated by, for example, determining a binding constant for the formation of the complex. The binding constant is the value obtained when the concentration of the complex is divided by the product of the component concentrations. In general, two compounds are said to "bind," in the context of the present invention, when the binding constant for complex formation exceeds about 10^3 L/mol. The binding constant may be determined using methods well known in the art.

Binding agents may be further capable of differentiating between patients with and without a cancer, such as colon cancer, using the representative assays provided herein. In other words, antibodies or other binding agents that bind to a colon tumor protein will generate a signal indicating the presence of a cancer in at least about 20% of patients with the disease, and will generate a negative signal indicating the absence of the disease in at least about 90% of individuals without the cancer. To determine whether a binding agent satisfies this requirement, biological samples (*e.g.*, blood, sera, sputum, urine and/or tumor biopsies) from patients with and without a cancer (as determined using standard clinical tests) may be assayed as described herein for the presence of polypeptides that bind to the binding agent. It will be apparent that a statistically significant number of samples with and without the disease should be assayed. Each binding agent should satisfy the above criteria; however, those of ordinary skill in the art will recognize that binding agents may be used in combination to improve sensitivity.

Any agent that satisfies the above requirements may be a binding agent. For example, a binding agent may be a ribosome, with or without a peptide component, an RNA molecule or a polypeptide. In a preferred embodiment, a binding agent is an

antibody or an antigen-binding fragment thereof. Antibodies may be prepared by any of a variety of techniques known to those of ordinary skill in the art. *See, e.g.,* Harlow and Lane, *Antibodies: A Laboratory Manual*, Cold Spring Harbor Laboratory, 1988. In general, antibodies can be produced by cell culture techniques, including the generation of monoclonal antibodies as described herein, or via transfection of antibody genes into suitable bacterial or mammalian cell hosts, in order to allow for the production of recombinant antibodies. In one technique, an immunogen comprising the polypeptide is initially injected into any of a wide variety of mammals (*e.g.,* mice, rats, rabbits, sheep or goats). In this step, the polypeptides of this invention may serve as the immunogen without modification. Alternatively, particularly for relatively short polypeptides, a superior immune response may be elicited if the polypeptide is joined to a carrier protein, such as bovine serum albumin or keyhole limpet hemocyanin. The immunogen is injected into the animal host, preferably according to a predetermined schedule incorporating one or more booster immunizations, and the animals are bled periodically. Polyclonal antibodies specific for the polypeptide may then be purified from such antisera by, for example, affinity chromatography using the polypeptide coupled to a suitable solid support.

Monoclonal antibodies specific for an antigenic polypeptide of interest may be prepared, for example, using the technique of Kohler and Milstein, *Eur. J. Immunol.* 6:511-519, 1976, and improvements thereto. Briefly, these methods involve the preparation of immortal cell lines capable of producing antibodies having the desired specificity (*i.e.,* reactivity with the polypeptide of interest). Such cell lines may be produced, for example, from spleen cells obtained from an animal immunized as described above. The spleen cells are then immortalized by, for example, fusion with a myeloma cell fusion partner, preferably one that is syngeneic with the immunized animal. A variety of fusion techniques may be employed. For example, the spleen cells and myeloma cells may be combined with a nonionic detergent for a few minutes and then plated at low density on a selective medium that supports the growth of hybrid cells, but not myeloma cells. A preferred selection technique uses HAT (hypoxanthine, aminopterin, thymidine) selection. After a sufficient time, usually about 1 to 2 weeks, colonies of hybrids are observed. Single colonies are selected and their culture

supernatants tested for binding activity against the polypeptide. Hybridomas having high reactivity and specificity are preferred.

Monoclonal antibodies may be isolated from the supernatants of growing hybridoma colonies. In addition, various techniques may be employed to enhance the yield, such as injection of the hybridoma cell line into the peritoneal cavity of a suitable vertebrate host, such as a mouse. Monoclonal antibodies may then be harvested from the ascites fluid or the blood. Contaminants may be removed from the antibodies by conventional techniques, such as chromatography, gel filtration, precipitation, and extraction. The polypeptides of this invention may be used in the purification process in, for example, an affinity chromatography step.

Within certain embodiments, the use of antigen-binding fragments of antibodies may be preferred. Such fragments include Fab fragments, which may be prepared using standard techniques. Briefly, immunoglobulins may be purified from rabbit serum by affinity chromatography on Protein A bead columns (Harlow and Lane, *Antibodies: A Laboratory Manual*, Cold Spring Harbor Laboratory, 1988) and digested by papain to yield Fab and Fc fragments. The Fab and Fc fragments may be separated by affinity chromatography on protein A bead columns.

Monoclonal antibodies of the present invention may be coupled to one or more therapeutic agents. Suitable agents in this regard include radionuclides, differentiation inducers, drugs, toxins, and derivatives thereof. Preferred radionuclides include ^{90}Y , ^{123}I , ^{125}I , ^{131}I , ^{186}Re , ^{188}Re , ^{211}At , and ^{213}Bi . Preferred drugs include methotrexate, and pyrimidine and purine analogs. Preferred differentiation inducers include phorbol esters and butyric acid. Preferred toxins include ricin, abrin, diphtheria toxin, cholera toxin, gelonin, *Pseudomonas* exotoxin, Shigella toxin, and pokeweed antiviral protein.

A therapeutic agent may be coupled (e.g., covalently bonded) to a suitable monoclonal antibody either directly or indirectly (e.g., via a linker group). A direct reaction between an agent and an antibody is possible when each possesses a substituent capable of reacting with the other. For example, a nucleophilic group, such as an amino or sulfhydryl group, on one may be capable of reacting with a carbonyl-

containing group, such as an anhydride or an acid halide, or with an alkyl group containing a good leaving group (e.g., a halide) on the other.

Alternatively, it may be desirable to couple a therapeutic agent and an antibody via a linker group. A linker group can function as a spacer to distance an antibody from an agent in order to avoid interference with binding capabilities. A linker group can also serve to increase the chemical reactivity of a substituent on an agent or an antibody, and thus increase the coupling efficiency. An increase in chemical reactivity may also facilitate the use of agents, or functional groups on agents, which otherwise would not be possible.

It will be evident to those skilled in the art that a variety of bifunctional or polyfunctional reagents, both homo- and hetero-functional (such as those described in the catalog of the Pierce Chemical Co., Rockford, IL), may be employed as the linker group. Coupling may be effected, for example, through amino groups, carboxyl groups, sulfhydryl groups or oxidized carbohydrate residues. There are numerous references describing such methodology, e.g., U.S. Patent No. 4,671,958, to Rodwell et al.

Where a therapeutic agent is more potent when free from the antibody portion of the immunoconjugates of the present invention, it may be desirable to use a linker group which is cleavable during or upon internalization into a cell. A number of different cleavable linker groups have been described. The mechanisms for the intracellular release of an agent from these linker groups include cleavage by reduction of a disulfide bond (e.g., U.S. Patent No. 4,489,710, to Spitler), by irradiation of a photolabile bond (e.g., U.S. Patent No. 4,625,014, to Senter et al.), by hydrolysis of derivatized amino acid side chains (e.g., U.S. Patent No. 4,638,045, to Kohn et al.), by serum complement-mediated hydrolysis (e.g., U.S. Patent No. 4,671,958, to Rodwell et al.), and acid-catalyzed hydrolysis (e.g., U.S. Patent No. 4,569,789, to Blattler et al.).

It may be desirable to couple more than one agent to an antibody. In one embodiment, multiple molecules of an agent are coupled to one antibody molecule. In another embodiment, more than one type of agent may be coupled to one antibody. Regardless of the particular embodiment, immunoconjugates with more than one agent may be prepared in a variety of ways. For example, more than one agent may be

coupled directly to an antibody molecule, or linkers that provide multiple sites for attachment can be used. Alternatively, a carrier can be used.

A carrier may bear the agents in a variety of ways, including covalent bonding either directly or via a linker group. Suitable carriers include proteins such as albumins (e.g., U.S. Patent No. 4,507,234, to Kato et al.), peptides and polysaccharides such as aminodextran (e.g., U.S. Patent No. 4,699,784, to Shih et al.). A carrier may also bear an agent by noncovalent bonding or by encapsulation, such as within a liposome vesicle (e.g., U.S. Patent Nos. 4,429,008 and 4,873,088). Carriers specific for radionuclide agents include radiohalogenated small molecules and chelating compounds. For example, U.S. Patent No. 4,735,792 discloses representative radiohalogenated small molecules and their synthesis. A radionuclide chelate may be formed from chelating compounds that include those containing nitrogen and sulfur atoms as the donor atoms for binding the metal, or metal oxide, radionuclide. For example, U.S. Patent No. 4,673,562, to Davison et al. discloses representative chelating compounds and their synthesis.

A variety of routes of administration for the antibodies and immunoconjugates may be used. Typically, administration will be intravenous, intramuscular, subcutaneous or in the bed of a resected tumor. It will be evident that the precise dose of the antibody/immunoconjugate will vary depending upon the antibody used, the antigen density on the tumor, and the rate of clearance of the antibody.

T CELLS

Immunotherapeutic compositions may also, or alternatively, comprise T cells specific for a colon tumor protein. Such cells may generally be prepared *in vitro* or *ex vivo*, using standard procedures. For example, T cells may be isolated from bone marrow, peripheral blood, or a fraction of bone marrow or peripheral blood of a patient, using a commercially available cell separation system, such as the Isolex™ System, available from Nexell Therapeutics, Inc. (Irvine, CA; see also U.S. Patent No. 5,240,856; U.S. Patent No. 5,215,926; WO 89/06280; WO 91/16116 and WO

92/07243). Alternatively, T cells may be derived from related or unrelated humans, non-human mammals, cell lines or cultures.

T cells may be stimulated with a colon tumor polypeptide, polynucleotide encoding a colon tumor polypeptide and/or an antigen presenting cell (APC) that expresses such a polypeptide. Such stimulation is performed under conditions and for a time sufficient to permit the generation of T cells that are specific for the polypeptide. Preferably, a colon tumor polypeptide or polynucleotide is present within a delivery vehicle, such as a microsphere, to facilitate the generation of specific T cells.

T cells are considered to be specific for a colon tumor polypeptide if the T cells specifically proliferate, secrete cytokines or kill target cells coated with the polypeptide or expressing a gene encoding the polypeptide. T cell specificity may be evaluated using any of a variety of standard techniques. For example, within a chromium release assay or proliferation assay, a stimulation index of more than two fold increase in lysis and/or proliferation, compared to negative controls, indicates T cell specificity. Such assays may be performed, for example, as described in Chen et al., *Cancer Res.* 54:1065-1070, 1994. Alternatively, detection of the proliferation of T cells may be accomplished by a variety of known techniques. For example, T cell proliferation can be detected by measuring an increased rate of DNA synthesis (e.g., by pulse-labeling cultures of T cells with tritiated thymidine and measuring the amount of tritiated thymidine incorporated into DNA). Contact with a colon tumor polypeptide (100 ng/ml - 100 µg/ml, preferably 200 ng/ml - 25 µg/ml) for 3 - 7 days should result in at least a two fold increase in proliferation of the T cells. Contact as described above for 2-3 hours should result in activation of the T cells, as measured using standard cytokine assays in which a two fold increase in the level of cytokine release (e.g., TNF or IFN-γ) is indicative of T cell activation (see Coligan et al., *Current Protocols in Immunology*, vol. 1, Wiley Interscience (Greene 1998)). T cells that have been activated in response to a colon tumor polypeptide, polynucleotide or polypeptide-expressing APC may be CD4⁺ and/or CD8⁺. Colon tumor protein-specific T cells may be expanded using standard techniques. Within preferred embodiments, the T cells are

derived from a patient, a related donor or an unrelated donor, and are administered to the patient following stimulation and expansion.

- For therapeutic purposes, CD4⁺ or CD8⁺ T cells that proliferate in response to a colon tumor polypeptide, polynucleotide or APC can be expanded in number either *in vitro* or *in vivo*. Proliferation of such T cells *in vitro* may be accomplished in a variety of ways. For example, the T cells can be re-exposed to a colon tumor polypeptide, or a short peptide corresponding to an immunogenic portion of such a polypeptide, with or without the addition of T cell growth factors, such as interleukin-2, and/or stimulator cells that synthesize a colon tumor polypeptide. Alternatively, one or more T cells that proliferate in the presence of a colon tumor protein can be expanded in number by cloning. Methods for cloning cells are well known in the art, and include limiting dilution.

PHARMACEUTICAL COMPOSITIONS AND VACCINES

- Within certain aspects, polypeptides, polynucleotides, T cells and/or binding agents described herein may be incorporated into pharmaceutical compositions or immunogenic compositions (*i.e.*, vaccines). Pharmaceutical compositions comprise one or more such compounds and a physiologically acceptable carrier. Vaccines may comprise one or more such compounds and an immunostimulant. An immunostimulant may be any substance that enhances or potentiates an immune response (antibody and/or cell-mediated) to an exogenous antigen. Examples of immunostimulants include adjuvants, biodegradable microspheres (*e.g.*, polylactic galactide) and liposomes (into which the compound is incorporated; *see e.g.*, Fullerton, U.S. Patent No. 4,235,877). Vaccine preparation is generally described in, for example, M.F. Powell and M.J. Newman, eds., "Vaccine Design (the subunit and adjuvant approach)," Plenum Press (NY, 1995). Pharmaceutical compositions and vaccines within the scope of the present invention may also contain other compounds, which may be biologically active or inactive. For example, one or more immunogenic portions of other tumor antigens may be present, either incorporated into a fusion polypeptide or as a separate compound, within the composition or vaccine.

- A pharmaceutical composition or vaccine may contain DNA encoding one or more of the polypeptides as described above, such that the polypeptide is generated *in situ*. As noted above, the DNA may be present within any of a variety of delivery systems known to those of ordinary skill in the art, including nucleic acid expression systems, bacteria and viral expression systems. Numerous gene delivery techniques are well known in the art, such as those described by Rolland, *Crit. Rev. Therap. Drug Carrier Systems* 15:143-198, 1998, and references cited therein. Appropriate nucleic acid expression systems contain the necessary DNA sequences for expression in the patient (such as a suitable promoter and terminating signal). Bacterial delivery systems involve the administration of a bacterium (such as *Bacillus-Calmette-Guerrin*) that expresses an immunogenic portion of the polypeptide on its cell surface or secretes such an epitope. In a preferred embodiment, the DNA may be introduced using a viral expression system (e.g., vaccinia or other pox virus, retrovirus, or adenovirus), which may involve the use of a non-pathogenic (defective), replication competent virus. Suitable systems are disclosed, for example, in Fisher-Hoch et al., *Proc. Natl. Acad. Sci. USA* 86:317-321, 1989; Flexner et al., *Ann. N.Y. Acad. Sci.* 569:86-103, 1989; Flexner et al., *Vaccine* 8:17-21, 1990; U.S. Patent Nos. 4,603,112, 4,769,330, and 5,017,487; WO 89/01973; U.S. Patent No. 4,777,127; GB 2,200,651; EP 0,345,242; WO 91/02805; Berkner, *Biotechniques* 6:616-627, 1988; Rosenfeld et al., *Science* 252:431-434, 1991; Kolls et al., *Proc. Natl. Acad. Sci. USA* 91:215-219, 1994; Kass-Eisler et al., *Proc. Natl. Acad. Sci. USA* 90:11498-11502, 1993; Guzman et al., *Circulation* 88:2838-2848, 1993; and Guzman et al., *Cir. Res.* 73:1202-1207, 1993. Techniques for incorporating DNA into such expression systems are well known to those of ordinary skill in the art. The DNA may also be "naked," as described, for example, in Ulmer et al., *Science* 259:1745-1749, 1993 and reviewed by Cohen, *Science* 259:1691-1692, 1993. The uptake of naked DNA may be increased by coating the DNA onto biodegradable beads, which are efficiently transported into the cells. It will be apparent that a vaccine may comprise both a polynucleotide and a polypeptide component. Such vaccines may provide for an enhanced immune response.
- It will be apparent that a vaccine may contain pharmaceutically acceptable salts of the polynucleotides and polypeptides provided herein. Such salts

may be prepared from pharmaceutically acceptable non-toxic bases, including organic bases (e.g., salts of primary, secondary and tertiary amines and basic amino acids) and inorganic bases (e.g., sodium, potassium, lithium, ammonium, calcium and magnesium salts).

5 While any suitable carrier known to those of ordinary skill in the art may be employed in the pharmaceutical compositions of this invention, the type of carrier will vary depending on the mode of administration. Compositions of the present invention may be formulated for any appropriate manner of administration, including for example, topical, oral, nasal, intravenous, intracranial, intraperitoneal, subcutaneous
10 or intramuscular administration. For parenteral administration, such as subcutaneous injection, the carrier preferably comprises water, saline, alcohol, a fat, a wax or a buffer. For oral administration, any of the above carriers or a solid carrier, such as mannitol, lactose, starch, magnesium stearate, sodium saccharine, talcum, cellulose, glucose, sucrose, and magnesium carbonate, may be employed. Biodegradable microspheres
15 (e.g., polylactate polyglycolate) may also be employed as carriers for the pharmaceutical compositions of this invention. Suitable biodegradable microspheres are disclosed, for example, in U.S. Patent Nos. 4,897,268; 5,075,109; 5,928,647; 5,811,128; 5,820,883; 5,853,763; 5,814,344 and 5,942,252.

Such compositions may also comprise buffers (e.g., neutral buffered
20 saline or phosphate buffered saline), carbohydrates (e.g., glucose, mannose, sucrose or dextrans), mannitol, proteins, polypeptides or amino acids such as glycine, antioxidants, bacteriostats, chelating agents such as EDTA or glutathione, adjuvants (e.g., aluminum hydroxide), solutes that render the formulation isotonic, hypotonic or weakly hypertonic with the blood of a recipient, suspending agents, thickening agents and/or preservatives.
25 Alternatively, compositions of the present invention may be formulated as a lyophilizate. Compounds may also be encapsulated within liposomes using well known technology.

Any of a variety of immunostimulants may be employed in the vaccines of this invention. For example, an adjuvant may be included. Most adjuvants contain a
30 substance designed to protect the antigen from rapid catabolism, such as aluminum hydroxide or mineral oil, and a stimulator of immune responses, such as lipid A,

Bordetella pertussis or *Mycobacterium tuberculosis* derived proteins. Suitable adjuvants are commercially available as, for example, Freund's Incomplete Adjuvant and Complete Adjuvant (Difco Laboratories, Detroit, MI); Merck Adjuvant 65 (Merck and Company, Inc., Rahway, NJ); AS-2 (SmithKline Beecham, Philadelphia, PA);
5 aluminum salts such as aluminum hydroxide gel (alum) or aluminum phosphate; salts of calcium, iron or zinc; an insoluble suspension of acylated tyrosine; acylated sugars; cationically or anionically derivatized polysaccharides; polyphosphazenes; biodegradable microspheres; monophosphoryl lipid A and quil A. Cytokines, such as GM-CSF or interleukin-2, -7, or -12, may also be used as adjuvants.

10 Within the vaccines provided herein, the adjuvant composition is preferably designed to induce an immune response predominantly of the Th1 type. High levels of Th1-type cytokines (e.g., IFN- γ , TNF α , IL-2 and IL-12) tend to favor the induction of cell mediated immune responses to an administered antigen. In contrast,
15 high levels of Th2-type cytokines (e.g., IL-4, IL-5, IL-6 and IL-10) tend to favor the induction of humoral immune responses. Following application of a vaccine as provided herein, a patient will support an immune response that includes Th1- and Th2-type responses. Within a preferred embodiment, in which a response is predominantly Th1-type, the level of Th1-type cytokines will increase to a greater extent than the level of Th2-type cytokines. The levels of these cytokines may be readily assessed using
20 standard assays. For a review of the families of cytokines, see Mosmann and Coffman, *Ann. Rev. Immunol.* 7:145-173, 1989.

Preferred adjuvants for use in eliciting a predominantly Th1-type response include, for example, a combination of monophosphoryl lipid A, preferably 3-de-O-acylated monophosphoryl lipid A (3D-MPL), together with an aluminum salt.
25 MPL adjuvants are available from Corixa Corporation (Seattle, WA; see US Patent Nos. 4,436,727; 4,877,611; 4,866,034 and 4,912,094). CpG-containing oligonucleotides (in which the CpG dinucleotide is unmethylated) also induce a predominantly Th1 response. Such oligonucleotides are well known and are described, for example, in WO 96/02555 and WO 99/33488. Immunostimulatory DNA sequences
30 are also described, for example, by Sato et al., *Science* 273:352, 1996. Another preferred adjuvant is a saponin, preferably QS21 (Aquila Biopharmaceuticals Inc.,

Framingham, MA), which may be used alone or in combination with other adjuvants. For example, an enhanced system involves the combination of a monophosphoryl lipid A and saponin derivative, such as the combination of QS21 and 3D-MPL as described in WO 94/00153, or a less reactogenic composition where the QS21 is quenched with cholesterol, as described in WO 96/33739. Other preferred formulations comprise an oil-in-water emulsion and tocopherol. A particularly potent adjuvant formulation involving QS21, 3D-MPL and tocopherol in an oil-in-water emulsion is described in WO 95/17210.

Other preferred adjuvants include Montanide ISA 720 (Seppic, France), SAF (Chiron, California, United States), ISCOMS (CSL), MF-59 (Chiron), the SBAS series of adjuvants (e.g., SBAS-2 or SBAS-4, available from SmithKline Beecham, Rixensart, Belgium), Detox (Ribi ImmunoChem Research Inc., Hamilton, MT), RC-529 (Ribi ImmunoChem Research Inc., Hamilton, MT) and Aminoalkyl glucosaminide 4-phosphates (AGPs).

Any vaccine provided herein may be prepared using well known methods that result in a combination of antigen, immune response enhancer and a suitable carrier or excipient. The compositions described herein may be administered as part of a sustained release formulation (i.e., a formulation such as a capsule, sponge or gel (composed of polysaccharides, for example) that effects a slow release of compound following administration). Such formulations may generally be prepared using well known technology (see, e.g., Coombes et al., *Vaccine 14*:1429-1438, 1996) and administered by, for example, oral, rectal or subcutaneous implantation, or by implantation at the desired target site. Sustained-release formulations may contain a polypeptide, polynucleotide or antibody dispersed in a carrier matrix and/or contained within a reservoir surrounded by a rate controlling membrane.

Carriers for use within such formulations are biocompatible, and may also be biodegradable; preferably the formulation provides a relatively constant level of active component release. Such carriers include microparticles of poly(lactide-co-glycolide), as well as polyacrylate, latex, starch, cellulose and dextran. Other delayed-release carriers include supramolecular biovectors, which comprise a non-liquid hydrophilic core (e.g., a cross-linked polysaccharide or oligosaccharide) and,

optionally, an external layer comprising an amphiphilic compound, such as a phospholipid (*see e.g.*, U.S. Patent No. 5,151,254 and PCT applications WO 94/20078, WO/94/23701 and WO 96/06638). The amount of active compound contained within a sustained release formulation depends upon the site of implantation, the rate and expected duration of release and the nature of the condition to be treated or prevented.

Any of a variety of delivery vehicles may be employed within pharmaceutical compositions and vaccines to facilitate production of an antigen-specific immune response that targets tumor cells. Delivery vehicles include antigen-presenting cells (APCs), such as dendritic cells, macrophages, B cells, monocytes and other cells that may be engineered to be efficient APCs. Such cells may, but need not, be genetically modified to increase the capacity for presenting the antigen, to improve activation and/or maintenance of the T cell response, to have anti-tumor effects *per se* and/or to be immunologically compatible with the receiver (*i.e.*, matched HLA haplotype). APCs may generally be isolated from any of a variety of biological fluids and organs, including tumor and peritumoral tissues, and may be autologous, allogeneic, syngeneic or xenogeneic cells.

Certain preferred embodiments of the present invention use dendritic cells or progenitors thereof as antigen-presenting cells. Dendritic cells are highly potent APCs (Banchereau and Steinman, *Nature* 392:245-251, 1998) and have been shown to be effective as a physiological adjuvant for eliciting prophylactic or therapeutic antitumor immunity (*see* Timmerman and Levy, *Ann. Rev. Med.* 50:507-529, 1999). In general, dendritic cells may be identified based on their typical shape (stellate *in situ*, with marked cytoplasmic processes (dendrites) visible *in vitro*), their ability to take up, process and present antigens with high efficiency and their ability to activate naïve T cell responses. Dendritic cells may, of course, be engineered to express specific cell-surface receptors or ligands that are not commonly found on dendritic cells *in vivo* or *ex vivo*, and such modified dendritic cells are contemplated by the present invention. As an alternative to dendritic cells, secreted vesicles antigen-loaded dendritic cells (called exosomes) may be used within a vaccine (*see* Zitvogel et al., *Nature Med.* 4:594-600, 1998).

Dendritic cells and progenitors may be obtained from peripheral blood, bone marrow, tumor-infiltrating cells, peritumoral tissues-infiltrating cells, lymph nodes, spleen, skin, umbilical cord blood or any other suitable tissue or fluid. For example, dendritic cells may be differentiated *ex vivo* by adding a combination of
5 cytokines such as GM-CSF, IL-4, IL-13 and/or TNF α to cultures of monocytes harvested from peripheral blood. Alternatively, CD34 positive cells harvested from peripheral blood, umbilical cord blood or bone marrow may be differentiated into dendritic cells by adding to the culture medium combinations of GM-CSF, IL-3, TNF α ,
10 CD40 ligand, LPS, flt3 ligand and/or other compound(s) that induce differentiation, maturation and proliferation of dendritic cells.

Dendritic cells are conveniently categorized as "immature" and "mature" cells, which allows a simple way to discriminate between two well characterized phenotypes. However, this nomenclature should not be construed to exclude all possible intermediate stages of differentiation. Immature dendritic cells are
15 characterized as APC with a high capacity for antigen uptake and processing, which correlates with the high expression of Fc γ receptor and mannose receptor. The mature phenotype is typically characterized by a lower expression of these markers, but a high expression of cell surface molecules responsible for T cell activation such as class I and class II MHC, adhesion molecules (*e.g.*, CD54 and CD11) and costimulatory molecules
20 (*e.g.*, CD40, CD80, CD86 and 4-1BB).

APCs may generally be transfected with a polynucleotide encoding a colon tumor protein (or portion or other variant thereof) such that the colon tumor polypeptide, or an immunogenic portion thereof, is expressed on the cell surface. Such transfection may take place *ex vivo*, and a composition or vaccine comprising such
25 transfected cells may then be used for therapeutic purposes, as described herein. Alternatively, a gene delivery vehicle that targets a dendritic or other antigen presenting cell may be administered to a patient, resulting in transfection that occurs *in vivo*. *In vivo* and *ex vivo* transfection of dendritic cells, for example, may generally be performed using any methods known in the art, such as those described in WO
30 97/24447, or the gene gun approach described by Mahvi et al., *Immunology and cell Biology* 75:456-460, 1997. Antigen loading of dendritic cells may be achieved by

- incubating dendritic cells or progenitor cells with the colon tumor polypeptide, DNA (naked or within a plasmid vector) or RNA; or with antigen-expressing recombinant bacterium or viruses (*e.g.*, vaccinia, fowlpox, adenovirus or lentivirus vectors). Prior to loading, the polypeptide may be covalently conjugated to an immunological partner that
- 5 provides T cell help (*e.g.*, a carrier molecule). Alternatively, a dendritic cell may be pulsed with a non-conjugated immunological partner, separately or in the presence of the polypeptide.

- Vaccines and pharmaceutical compositions may be presented in unit-dose or multi-dose containers, such as sealed ampoules or vials. Such containers are
- 10 preferably hermetically sealed to preserve sterility of the formulation until use. In general, formulations may be stored as suspensions, solutions or emulsions in oily or aqueous vehicles. Alternatively, a vaccine or pharmaceutical composition may be stored in a freeze-dried condition requiring only the addition of a sterile liquid carrier immediately prior to use.

15 CANCER THERAPY

- In further aspects of the present invention, the compositions described herein may be used for immunotherapy of cancer, such as colon cancer. Within such methods, pharmaceutical compositions and vaccines are typically administered to a patient. As used herein, a "patient" refers to any warm-blooded animal, preferably a
- 20 human. A patient may or may not be afflicted with cancer. Accordingly, the above pharmaceutical compositions and vaccines may be used to prevent the development of a cancer or to treat a patient afflicted with a cancer. A cancer may be diagnosed using criteria generally accepted in the art, including the presence of a malignant tumor. Pharmaceutical compositions and vaccines may be administered either prior to or
- 25 following surgical removal of primary tumors and/or treatment such as administration of radiotherapy or conventional chemotherapeutic drugs. Administration may be by any suitable method, including administration by intravenous, intraperitoneal, intramuscular, subcutaneous, intranasal, intradermal, anal, vaginal, topical and oral routes.

Within certain embodiments, immunotherapy may be active immunotherapy, in which treatment relies on the *in vivo* stimulation of the endogenous host immune system to react against tumors with the administration of immune response-modifying agents (such as polypeptides and polynucleotides as provided
5 herein).

Within other embodiments, immunotherapy may be passive immunotherapy, in which treatment involves the delivery of agents with established tumor-immune reactivity (such as effector cells or antibodies) that can directly or indirectly mediate antitumor effects and does not necessarily depend on an intact host
10 immune system. Examples of effector cells include T cells as discussed above, T lymphocytes (such as CD8⁺ cytotoxic T lymphocytes and CD4⁺ T-helper tumor-infiltrating lymphocytes), killer cells (such as Natural Killer cells and lymphokine-activated killer cells), B cells and antigen-presenting cells (such as dendritic cells and macrophages) expressing a polypeptide provided herein. T cell receptors and antibody
15 receptors specific for the polypeptides recited herein may be cloned, expressed and transferred into other vectors or effector cells for adoptive immunotherapy. The polypeptides provided herein may also be used to generate antibodies or anti-idiotypic antibodies (as described above and in U.S. Patent No. 4,918,164) for passive immunotherapy.

20 Effector cells may generally be obtained in sufficient quantities for adoptive immunotherapy by growth *in vitro*, as described herein. Culture conditions for expanding single antigen-specific effector cells to several billion in number with retention of antigen recognition *in vivo* are well known in the art. Such *in vitro* culture conditions typically use intermittent stimulation with antigen, often in the presence of
25 cytokines (such as IL-2) and non-dividing feeder cells. As noted above, immunoreactive polypeptides as provided herein may be used to rapidly expand antigen-specific T cell cultures in order to generate a sufficient number of cells for immunotherapy. In particular, antigen-presenting cells, such as dendritic, macrophage, monocyte, fibroblast and/or B cells, may be pulsed with immunoreactive polypeptides
30 or transfected with one or more polynucleotides using standard techniques well known in the art. For example, antigen-presenting cells can be transfected with a

polynucleotide having a promoter appropriate for increasing expression in a recombinant virus or other expression system. Cultured effector cells for use in therapy must be able to grow and distribute widely, and to survive long term *in vivo*. Studies have shown that cultured effector cells can be induced to grow *in vivo* and to survive
5 long term in substantial numbers by repeated stimulation with antigen supplemented with IL-2 (see, for example, Cheever et al., *Immunological Reviews* 157:177, 1997).

Alternatively, a vector expressing a polypeptide recited herein may be introduced into antigen presenting cells taken from a patient and clonally propagated *ex vivo* for transplant back into the same patient. Transfected cells may be reintroduced
10 into the patient using any means known in the art, preferably in sterile form by intravenous, intracavitary, intraperitoneal or intratumor administration.

Routes and frequency of administration of the therapeutic compositions described herein, as well as dosage, will vary from individual to individual, and may be readily established using standard techniques. In general, the pharmaceutical
15 compositions and vaccines may be administered by injection (e.g., intracutaneous, intramuscular, intravenous or subcutaneous), intranasally (e.g., by aspiration) or orally. Preferably, between 1 and 10 doses may be administered over a 52 week period. Preferably, 6 doses are administered, at intervals of 1 month, and booster vaccinations may be given periodically thereafter. Alternate protocols may be appropriate for
20 individual patients. A suitable dose is an amount of a compound that, when administered as described above, is capable of promoting an anti-tumor immune response, and is at least 10-50% above the basal (i.e., untreated) level. Such response can be monitored by measuring the anti-tumor antibodies in a patient or by vaccine-dependent generation of cytolytic effector cells capable of killing the patient's tumor
25 cells *in vitro*. Such vaccines should also be capable of causing an immune response that leads to an improved clinical outcome (e.g., more frequent remissions, complete or partial or longer disease-free survival) in vaccinated patients as compared to non-vaccinated patients. In general, for pharmaceutical compositions and vaccines comprising one or more polypeptides, the amount of each polypeptide present in a dose
30 ranges from about 25 µg to 5 mg per kg of host. Suitable dose sizes will vary with the size of the patient, but will typically range from about 0.1 mL to about 5 mL.

In general, an appropriate dosage and treatment regimen provides the active compound(s) in an amount sufficient to provide therapeutic and/or prophylactic benefit. Such a response can be monitored by establishing an improved clinical outcome (e.g., more frequent remissions, complete or partial, or longer disease-free survival) in treated patients as compared to non-treated patients. Increases in preexisting immune responses to a colon tumor protein generally correlate with an improved clinical outcome. Such immune responses may generally be evaluated using standard proliferation, cytotoxicity or cytokine assays, which may be performed using samples obtained from a patient before and after treatment.

10 METHODS FOR DETECTING CANCER

In general, a cancer may be detected in a patient based on the presence of one or more colon tumor proteins and/or polynucleotides encoding such proteins in a biological sample (for example, blood, sera, sputum urine and/or tumor biopsies) obtained from the patient. In other words, such proteins may be used as markers to indicate the presence or absence of a cancer such as colon cancer. In addition, such proteins may be useful for the detection of other cancers. The binding agents provided herein generally permit detection of the level of antigen that binds to the agent in the biological sample. Polynucleotide primers and probes may be used to detect the level of mRNA encoding a tumor protein, which is also indicative of the presence or absence of a cancer. In general, a colon tumor sequence should be present at a level that is at least three fold higher in tumor tissue than in normal tissue

There are a variety of assay formats known to those of ordinary skill in the art for using a binding agent to detect polypeptide markers in a sample. See, e.g., Harlow and Lane, *Antibodies: A Laboratory Manual*, Cold Spring Harbor Laboratory, 1988. In general, the presence or absence of a cancer in a patient may be determined by (a) contacting a biological sample obtained from a patient with a binding agent; (b) detecting in the sample a level of polypeptide that binds to the binding agent; and (c) comparing the level of polypeptide with a predetermined cut-off value.

In a preferred embodiment, the assay involves the use of binding agent immobilized on a solid support to bind to and remove the polypeptide from the

remainder of the sample. The bound polypeptide may then be detected using a detection reagent that contains a reporter group and specifically binds to the binding agent/polypeptide complex. Such detection reagents may comprise, for example, a binding agent that specifically binds to the polypeptide or an antibody or other agent that specifically binds to the binding agent, such as an anti-immunoglobulin, protein G, protein A or a lectin. Alternatively, a competitive assay may be utilized, in which a polypeptide is labeled with a reporter group and allowed to bind to the immobilized binding agent after incubation of the binding agent with the sample. The extent to which components of the sample inhibit the binding of the labeled polypeptide to the binding agent is indicative of the reactivity of the sample with the immobilized binding agent. Suitable polypeptides for use within such assays include full length colon tumor proteins and portions thereof to which the binding agent binds, as described above.

The solid support may be any material known to those of ordinary skill in the art to which the tumor protein may be attached. For example, the solid support may be a test well in a microtiter plate or a nitrocellulose or other suitable membrane. Alternatively, the support may be a bead or disc, such as glass, fiberglass, latex or a plastic material such as polystyrene or polyvinylchloride. The support may also be a magnetic particle or a fiber optic sensor, such as those disclosed, for example, in U.S. Patent No. 5,359,681. The binding agent may be immobilized on the solid support using a variety of techniques known to those of skill in the art, which are amply described in the patent and scientific literature. In the context of the present invention, the term "immobilization" refers to both noncovalent association, such as adsorption, and covalent attachment (which may be a direct linkage between the agent and functional groups on the support or may be a linkage by way of a cross-linking agent). Immobilization by adsorption to a well in a microtiter plate or to a membrane is preferred. In such cases, adsorption may be achieved by contacting the binding agent, in a suitable buffer, with the solid support for a suitable amount of time. The contact time varies with temperature, but is typically between about 1 hour and about 1 day. In general, contacting a well of a plastic microtiter plate (such as polystyrene or polyvinylchloride) with an amount of binding agent ranging from about 10 ng to about

10 μg , and preferably about 100 ng to about 1 μg , is sufficient to immobilize an adequate amount of binding agent.

Covalent attachment of binding agent to a solid support may generally be achieved by first reacting the support with a bifunctional reagent that will react with
5 both the support and a functional group, such as a hydroxyl or amino group, on the binding agent. For example, the binding agent may be covalently attached to supports having an appropriate polymer coating using benzoquinone or by condensation of an aldehyde group on the support with an amine and an active hydrogen on the binding partner (*see, e.g.*, Pierce Immunotechnology Catalog and Handbook, 1991, at
10 A12-A13).

In certain embodiments, the assay is a two-antibody sandwich assay. This assay may be performed by first contacting an antibody that has been immobilized on a solid support, commonly the well of a microtiter plate, with the sample, such that polypeptides within the sample are allowed to bind to the immobilized antibody.
15 Unbound sample is then removed from the immobilized polypeptide-antibody complexes and a detection reagent (preferably a second antibody capable of binding to a different site on the polypeptide) containing a reporter group is added. The amount of detection reagent that remains bound to the solid support is then determined using a method appropriate for the specific reporter group.

20 More specifically, once the antibody is immobilized on the support as described above, the remaining protein binding sites on the support are typically blocked. Any suitable blocking agent known to those of ordinary skill in the art, such as bovine serum albumin or Tween 20™ (Sigma Chemical Co., St. Louis, MO). The immobilized antibody is then incubated with the sample, and polypeptide is allowed to
25 bind to the antibody. The sample may be diluted with a suitable diluent, such as phosphate-buffered saline (PBS) prior to incubation. In general, an appropriate contact time (*i.e.*, incubation time) is a period of time that is sufficient to detect the presence of polypeptide within a sample obtained from an individual with colon cancer. Preferably, the contact time is sufficient to achieve a level of binding that is at least about 95% of
30 that achieved at equilibrium between bound and unbound polypeptide. Those of ordinary skill in the art will recognize that the time necessary to achieve equilibrium

may be readily determined by assaying the level of binding that occurs over a period of time. At room temperature, an incubation time of about 30 minutes is generally sufficient.

Unbound sample may then be removed by washing the solid support
5 with an appropriate buffer, such as PBS containing 0.1% Tween 20™. The second antibody, which contains a reporter group, may then be added to the solid support. Preferred reporter groups include those groups recited above.

The detection reagent is then incubated with the immobilized antibody-polypeptide complex for an amount of time sufficient to detect the bound polypeptide.
10 An appropriate amount of time may generally be determined by assaying the level of binding that occurs over a period of time. Unbound detection reagent is then removed and bound detection reagent is detected using the reporter group. The method employed for detecting the reporter group depends upon the nature of the reporter group. For radioactive groups, scintillation counting or autoradiographic methods are
15 generally appropriate. Spectroscopic methods may be used to detect dyes, luminescent groups and fluorescent groups. Biotin may be detected using avidin, coupled to a different reporter group (commonly a radioactive or fluorescent group or an enzyme). Enzyme reporter groups may generally be detected by the addition of substrate (generally for a specific period of time), followed by spectroscopic or other analysis of
20 the reaction products.

To determine the presence or absence of a cancer, such as colon cancer, the signal detected from the reporter group that remains bound to the solid support is generally compared to a signal that corresponds to a predetermined cut-off value. In one preferred embodiment, the cut-off value for the detection of a cancer is the average
25 mean signal obtained when the immobilized antibody is incubated with samples from patients without the cancer. In general, a sample generating a signal that is three standard deviations above the predetermined cut-off value is considered positive for the cancer. In an alternate preferred embodiment, the cut-off value is determined using a Receiver Operator Curve, according to the method of Sackett et al., *Clinical*
30 *Epidemiology: A Basic Science for Clinical Medicine*, Little Brown and Co., 1985, p. 106-7. Briefly, in this embodiment, the cut-off value may be determined from a plot

of pairs of true positive rates (*i.e.*, sensitivity) and false positive rates (100%-specificity) that correspond to each possible cut-off value for the diagnostic test result. The cut-off value on the plot that is the closest to the upper left-hand corner (*i.e.*, the value that encloses the largest area) is the most accurate cut-off value, and a sample
5 generating a signal that is higher than the cut-off value determined by this method may be considered positive. Alternatively, the cut-off value may be shifted to the left along the plot, to minimize the false positive rate, or to the right, to minimize the false negative rate. In general, a sample generating a signal that is higher than the cut-off value determined by this method is considered positive for a cancer.

10 In a related embodiment, the assay is performed in a flow-through or strip test format, wherein the binding agent is immobilized on a membrane, such as nitrocellulose. In the flow-through test, polypeptides within the sample bind to the immobilized binding agent as the sample passes through the membrane. A second, labeled binding agent then binds to the binding agent-polypeptide complex as a solution
15 containing the second binding agent flows through the membrane. The detection of bound second binding agent may then be performed as described above. In the strip test format, one end of the membrane to which binding agent is bound is immersed in a solution containing the sample. The sample migrates along the membrane through a region containing second binding agent and to the area of immobilized binding agent.

20 Concentration of second binding agent at the area of immobilized antibody indicates the presence of a cancer. Typically, the concentration of second binding agent at that site generates a pattern, such as a line, that can be read visually. The absence of such a pattern indicates a negative result. In general, the amount of binding agent immobilized on the membrane is selected to generate a visually discernible pattern when the
25 biological sample contains a level of polypeptide that would be sufficient to generate a positive signal in the two-antibody sandwich assay, in the format discussed above. Preferred binding agents for use in such assays are antibodies and antigen-binding fragments thereof. Preferably, the amount of antibody immobilized on the membrane ranges from about 25 ng to about 1 μ g, and more preferably from about 50 ng to about
30 500 ng. Such tests can typically be performed with a very small amount of biological sample.

Of course, numerous other assay protocols exist that are suitable for use with the tumor proteins or binding agents of the present invention. The above descriptions are intended to be exemplary only. For example, it will be apparent to those of ordinary skill in the art that the above protocols may be readily modified to use
5 colon tumor polypeptides to detect antibodies that bind to such polypeptides in a biological sample. The detection of such colon tumor protein specific antibodies may correlate with the presence of a cancer.

A cancer may also, or alternatively, be detected based on the presence of T cells that specifically react with a colon tumor protein in a biological sample. Within
10 certain methods, a biological sample comprising $CD4^{+}$ and/or $CD8^{+}$ T cells isolated from a patient is incubated with a colon tumor polypeptide, a polynucleotide encoding such a polypeptide and/or an APC that expresses at least an immunogenic portion of such a polypeptide, and the presence or absence of specific activation of the T cells is detected. Suitable biological samples include, but are not limited to, isolated T cells.
15 For example, T cells may be isolated from a patient by routine techniques (such as by Ficoll/Hypaque density gradient centrifugation of peripheral blood lymphocytes). T cells may be incubated *in vitro* for 2-9 days (typically 4 days) at 37°C with polypeptide (e.g., 5 - 25 µg/ml). It may be desirable to incubate another aliquot of a T cell sample in the absence of colon tumor polypeptide to serve as a control. For $CD4^{+}$ T cells,
20 activation is preferably detected by evaluating proliferation of the T cells. For $CD8^{+}$ T cells, activation is preferably detected by evaluating cytolytic activity. A level of proliferation that is at least two fold greater and/or a level of cytolytic activity that is at least 20% greater than in disease-free patients indicates the presence of a cancer in the patient.

As noted above, a cancer may also, or alternatively, be detected based on the level of mRNA encoding a colon tumor protein in a biological sample. For example, at least two oligonucleotide primers may be employed in a polymerase chain reaction (PCR) based assay to amplify a portion of a colon tumor cDNA derived from a biological sample, wherein at least one of the oligonucleotide primers is specific for
30 (i.e., hybridizes to) a polynucleotide encoding the colon tumor protein. The amplified cDNA is then separated and detected using techniques well known in the art, such as

gel electrophoresis. Similarly, oligonucleotide probes that specifically hybridize to a polynucleotide encoding a colon tumor protein may be used in a hybridization assay to detect the presence of polynucleotide encoding the tumor protein in a biological sample.

To permit hybridization under assay conditions, oligonucleotide primers
5 and probes should comprise an oligonucleotide sequence that has at least about 60%, preferably at least about 75% and more preferably at least about 90%, identity to a portion of a polynucleotide encoding a colon tumor protein that is at least 10 nucleotides, and preferably at least 20 nucleotides, in length. Preferably,
10 oligonucleotide primers and/or probes hybridize to a polynucleotide encoding a polypeptide described herein under moderately stringent conditions, as defined above. Oligonucleotide primers and/or probes which may be usefully employed in the diagnostic methods described herein preferably are at least 10-40 nucleotides in length. In a preferred embodiment, the oligonucleotide primers comprise at least 10 contiguous nucleotides, more preferably at least 15 contiguous nucleotides, of a DNA molecule
15 having a sequence recited in SEQ ID NOs:1-1556. Techniques for both PCR based assays and hybridization assays are well known in the art (*see*, for example, Mullis et al., *Cold Spring Harbor Symp. Quant. Biol.*, 51:263, 1987; Erlich ed., *PCR Technology*, Stockton Press, NY, 1989).

One preferred assay employs RT-PCR, in which PCR is applied in
20 conjunction with reverse transcription. Typically, RNA is extracted from a biological sample, such as biopsy tissue, and is reverse transcribed to produce cDNA molecules. PCR amplification using at least one specific primer generates a cDNA molecule, which may be separated and visualized using, for example, gel electrophoresis. Amplification may be performed on biological samples taken from a test patient and
25 from an individual who is not afflicted with a cancer. The amplification reaction may be performed on several dilutions of cDNA spanning two orders of magnitude. A two-fold or greater increase in expression in several dilutions of the test patient sample as compared to the same dilutions of the non-cancerous sample is typically considered positive.

30 In another embodiment, the compositions described herein may be used as markers for the progression of cancer. In this embodiment, assays as described

above for the diagnosis of a cancer may be performed over time, and the change in the level of reactive polypeptide(s) or polynucleotide(s) evaluated. For example, the assays may be performed every 24-72 hours for a period of 6 months to 1 year, and thereafter performed as needed. In general, a cancer is progressing in those patients in whom the level of polypeptide or polynucleotide detected increases over time. In contrast, the cancer is not progressing when the level of reactive polypeptide or polynucleotide either remains constant or decreases with time.

Certain *in vivo* diagnostic assays may be performed directly on a tumor. One such assay involves contacting tumor cells with a binding agent. The bound binding agent may then be detected directly or indirectly via a reporter group. Such binding agents may also be used in histological applications. Alternatively, polynucleotide probes may be used within such applications.

As noted above, to improve sensitivity, multiple colon tumor protein markers may be assayed within a given sample. It will be apparent that binding agents specific for different proteins provided herein may be combined within a single assay. Further, multiple primers or probes may be used concurrently. The selection of tumor protein markers may be based on routine experiments to determine combinations that results in optimal sensitivity. In addition, or alternatively, assays for tumor proteins provided herein may be combined with assays for other known tumor antigens.

20 DIAGNOSTIC KITS

The present invention further provides kits for use within any of the above diagnostic methods. Such kits typically comprise two or more components necessary for performing a diagnostic assay. Components may be compounds, reagents, containers and/or equipment. For example, one container within a kit may contain a monoclonal antibody or fragment thereof that specifically binds to a colon tumor protein. Such antibodies or fragments may be provided attached to a support material, as described above. One or more additional containers may enclose elements, such as reagents or buffers, to be used in the assay. Such kits may also, or alternatively, contain a detection reagent as described above that contains a reporter group suitable for direct or indirect detection of antibody binding.

Alternatively, a kit may be designed to detect the level of mRNA encoding a colon tumor protein in a biological sample. Such kits generally comprise at least one oligonucleotide probe or primer, as described above, that hybridizes to a polynucleotide encoding a colon tumor protein. Such an oligonucleotide may be used, 5 for example, within a PCR or hybridization assay. Additional components that may be present within such kits include a second oligonucleotide and/or a diagnostic reagent or container to facilitate the detection of a polynucleotide encoding a colon tumor protein.

The following Examples are offered by way of illustration and not by way of limitation.

EXAMPLE 1

IDENTIFICATION OF COLON TUMOR PROTEIN CDNAS

This Example illustrates the identification of cDNA molecules encoding
5 colon tumor proteins using PCR-based cDNA subtraction methodology.

A pool of tester mRNA was collected from three colon adenocarcinoma
samples showing moderate histological differentiation and no evidence of metastasis.
Eight normal tissues, including brain, pancreas, bone marrow, liver, heart, lung,
stomach and small intestine were represented in the driver mRNA pool. cDNA
10 synthesis, hybridization and PCR amplification were performed according to the
methods of Clontech (Palo Alto, CA), with minor modifications. In a first subtraction,
the restriction enzymes PvuII, DraI, MscI and StuI were used to digest cDNAs. The
tester to driver ratio was 1:40. In a second subtraction, DraI, MscI and StuI were used
for cDNA digestion. A tester to driver ratio of 1:76 was employed. Following the PCR
15 amplification steps, the cDNAs were cloned into the pCR2.1 plasmid vector. The
libraries resulting from the first and second subtractions, named CPS1 and CPS2,
respectively, were used to obtain clones for microarray analysis and sequencing. Inserts
were PCR amplified and purified. Each clone was sequenced from one direction with
either M13 Forward primer or M13 Reverse primer. The determined cDNA sequences
20 for 1535 of the isolated clones are provided in SEQ ID NOs:1-1556.

A cDNA library was constructed in the PCR2.1 vector (Invitrogen,
Carlsbad, CA) by subtracting a pool of three colon tumors with a pool of normal colon,
spleen, brain, liver, kidney, lung, stomach and small intestine using PCR subtraction
methodologies (Clontech, Palo Alto, CA). The subtraction was performed using a
25 PCR-based protocol, which was modified to generate larger fragments. Within this
protocol, tester and driver double stranded cDNA were separately digested with five
restriction enzymes that recognize six-nucleotide restriction sites (MluI, MscI, PvuII,
SalI and StuI). This digestion resulted in an average cDNA size of 600 bp, rather than
the average size of 300 bp that results from digestion with RsaI according to the
30 Clontech protocol. This modification did not affect the subtraction efficiency. Two

tester populations were then created with different adapters, and the driver library remained without adapters.

The tester and driver libraries were then hybridized using excess driver cDNA. In the first hybridization step, driver was separately hybridized with each of the
5 two tester cDNA populations. This resulted in populations of (a) unhybridized tester cDNAs, (b) tester cDNAs hybridized to other tester cDNAs, (c) tester cDNAs hybridized to driver cDNAs, and (d) unhybridized driver cDNAs. The two separate hybridization reactions were then combined, and rehybridized in the presence of additional denatured driver cDNA. Following this second hybridization, in addition to
10 populations (a) through (d), a fifth population (e) was generated in which tester cDNA with one adapter hybridized to tester cDNA with the second adapter. Accordingly, the second hybridization step resulted in enrichment of differentially expressed sequences which could be used as templates for PCR amplification with adaptor-specific primers.

The ends were then filled in, and PCR amplification was performed
15 using adaptor-specific primers. Only population (e), which contained tester cDNA that did not hybridize to driver cDNA, was amplified exponentially. A second PCR amplification step was then performed, to reduce background and further enrich differentially expressed sequences.

This PCR-based subtraction technique normalizes differentially
20 expressed cDNAs so that rare transcripts that are over-expressed in colon tumor tissue may be recoverable. Such transcripts would be difficult to recover by traditional subtraction methods.

To characterize the complexity and redundancy of the subtracted library, 96 clones were randomly picked and 65 were sequenced, as previously described.
25 These sequences were further characterized by comparison with the most recent Genbank database (April, 1998) to determine their degree of novelty. No significant homologies were found to 21 of these clones, hereinafter referred to as 11092, 11093, 11096, 11098, 11103, 11174, 11108, 11112, 11115, 11117, 11118, 11134, 11151, 11154, 11158, 11168, 11172, 11175, 11184, 11185 and 11187. The determined cDNA
30 sequences for these clones are provided in SEQ ID NO: 48, 49, 52, 54, 59, 60, 65-69, 79, 89, 90, 93, 99-101 and 109-111, respectively.

- Two-thousand clones from the above mentioned cDNA subtraction library were randomly picked and submitted to a round of PCR amplification. Briefly, 0.5 µl of glycerol stock solution was added to 99.5 µl of pcr MIX (80 µl H₂O, 10 µl 10X PCR Buffer, 6 µl 25 mM MgCl₂, 1 µl 10 mM dNTPs, 1 µl 100 mM M13 forward primer (CACGACGTTGTAAAACGACGG), 1 µl 100 mM M13 reverse primer (CACAGGAAACAGCTATGACC)), and 0.5 µl 5 u/ml Taq polymerase (primers provided by (Operon Technologies, Alameda, CA). The PCR amplification was run for thirty cycles under the following conditions: 95°C for 5 min., 92°C for 30 sec., 57°C for 40 sec., 75°C for 2 min. and 75°C for 5 minutes.
- mRNA expression levels for representative clones were determined using microarray technology (Synteni, Palo Alto, CA) in colon tumor tissues (n=25), normal colon tissues (n=6), kidney, lung, liver, brain, heart, esophagus, small intestine, stomach, pancreas, adrenal gland, salivary gland, resting PBMC, activated PBMC, bone marrow, dendritic cells, spinal cord, blood vessels, skeletal muscle, skin, breast and fetal tissues. The number of tissue samples tested in each case was one (n=1), except where specifically noted above; additionally, all the above-mentioned tissues were derived from humans. The PCR amplification products were dotted onto slides in an array format, with each product occupying a unique location in the array. mRNA was extracted from the tissue sample to be tested, and fluorescent-labeled cDNA probes were generated by reverse transcription according to the protocol provided by Synteni. The microarrays were probed with the labeled cDNA probes, the slides scanned, and fluorescence intensity was measured. This intensity correlates with the hybridization intensity.
- Clones corresponding to SEQ ID Nos:1506-1556 were overexpressed in colon tumors and showed low or no expression levels in normal tissues.

EXAMPLE 2

SYNTHESIS OF POLYPEPTIDES

- Polypeptides may be synthesized on a Perkin Elmer/Applied Biosystems Division 430A peptide synthesizer using Fmoc chemistry with HPTU (O-

Benzotriazole-N,N',N'',N'''-tetramethyluronium hexafluorophosphate) activation. A Gly-Cys-Gly sequence may be attached to the amino terminus of the peptide to provide a method of conjugation, binding to an immobilized surface, or labeling of the peptide. Cleavage of the peptides from the solid support may be carried out using the following

5 cleavage mixture: trifluoroacetic acid:ethanedithiol:thioanisole:water:phenol (40:1:2:2:3). After cleaving for 2 hours, the peptides may be precipitated in cold methyl-t-butyl-ether. The peptide pellets may then be dissolved in water containing 0.1% trifluoroacetic acid (TFA) and lyophilized prior to purification by C18 reverse

10 phase HPLC. A gradient of 0%-60% acetonitrile (containing 0.1% TFA) in water (containing 0.1% TFA) may be used to elute the peptides. Following lyophilization of the pure fractions, the peptides may be characterized using electrospray or other types of mass spectrometry and by amino acid analysis.

From the foregoing it will be appreciated that, although specific embodiments of the invention have been described herein for purposes of illustration,

15 various modifications may be made without deviating from the spirit and scope of the invention. Accordingly, the invention is not limited except as by the appended claims.

CLAIMS

1. An isolated polypeptide, comprising at least an immunogenic portion of a colon tumor protein, or a variant thereof, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide sequence selected from the group consisting of:

- (a) sequences recited in SEQ ID NOs:1-1556;
- (b) sequences that hybridize to a sequence recited in any one of SEQ ID NOs:1-1556 under moderately stringent conditions; and
- (c) complements of sequences of (a) or (b).

2. An isolated polypeptide according to claim 1, wherein the polypeptide comprises an amino acid sequence that is encoded by a polynucleotide sequence recited in any one of SEQ ID NOs:1-1556 or a complement of any of the foregoing polynucleotide sequences.

3. An isolated polynucleotide encoding at least 15 amino acid residues of a colon tumor protein, or a variant thereof that differs in one or more substitutions, deletions, additions and/or insertions such that the ability of the variant to react with antigen-specific antisera is not substantially diminished, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide comprising a sequence recited in any one of SEQ ID NOs:1-1556 or a complement of any of the foregoing sequences.

4. An isolated polynucleotide encoding a colon tumor protein, or a variant thereof, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide comprising a sequence recited in any one of SEQ ID NOs:1-1556 or a complement of any of the foregoing sequences.

5. An isolated polynucleotide, comprising a sequence recited in any one of SEQ ID NOs:1-1556.

6. An isolated polynucleotide, comprising a sequence that hybridizes to a sequence recited in any one of SEQ ID NOs:1-1556 under moderately stringent conditions.

7. An isolated polynucleotide complementary to a polynucleotide according to any one of claims 3-6.

8. An expression vector, comprising a polynucleotide according to any one of claims 3-7.

9. A host cell transformed or transfected with an expression vector according to claim 8.

10. An isolated antibody, or antigen-binding fragment thereof, that specifically binds to a colon tumor protein that comprises an amino acid sequence that is encoded by a polynucleotide sequence recited in any one of SEQ ID NOs1-1556 or a complement of any of the foregoing polynucleotide sequences.

11. A fusion protein, comprising at least one polypeptide according to claim 1.

12. A fusion protein according to claim 11, wherein the fusion protein comprises an expression enhancer that increases expression of the fusion protein in a host cell transfected with a polynucleotide encoding the fusion protein.

13. A fusion protein according to claim 11, wherein the fusion protein comprises a T helper epitope that is not present within the polypeptide of claim 1.

14. A fusion protein according to claim 11, wherein the fusion protein comprises an affinity tag.

15. An isolated polynucleotide encoding a fusion protein according to claim 11.

16. A pharmaceutical composition, comprising a physiologically acceptable carrier and at least one component selected from the group consisting of:

- (a) a polypeptide according to claim 1;
- (b) a polynucleotide according to claim 3;
- (c) an antibody according to claim 10;
- (d) a fusion protein according to claim 11; and
- (e) a polynucleotide according to claim 15.

17. A vaccine comprising an immunostimulant and at least one component selected from the group consisting of:

- (a) a polypeptide according to claim 1;
- (b) a polynucleotide according to claim 3;
- (c) an antibody according to claim 10;
- (d) a fusion protein according to claim 11; and
- (e) a polynucleotide according to claim 15.

18. A vaccine according to claim 17, wherein the immunostimulant is an adjuvant.

19. A vaccine according to any claim 17, wherein the immunostimulant induces a predominantly Type I response.

20. A method for inhibiting the development of a cancer in a patient, comprising administering to a patient an effective amount of a pharmaceutical composition according to claim 16.

21. A method for inhibiting the development of a cancer in a patient, comprising administering to a patient an effective amount of a vaccine according to claim 17.

22. A pharmaceutical composition comprising an antigen-presenting cell that expresses a polypeptide according to claim 1, in combination with a pharmaceutically acceptable carrier or excipient.

23. A pharmaceutical composition according to claim 22, wherein the antigen presenting cell is a dendritic cell or a macrophage.

24. A vaccine comprising an antigen-presenting cell that expresses a polypeptide comprising at least an immunogenic portion of a colon tumor protein, or a variant thereof, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide sequence selected from the group consisting of:

- (a) sequences recited in SEQ ID NOs:1-1556;
 - (b) sequences that hybridize to a sequence recited in any one of SEQ ID NOs:1-1556 under moderately stringent conditions; and
 - (c) complements of sequences of (i) or (ii);
- in combination with an immunostimulant.

25. A vaccine according to claim 24, wherein the immunostimulant is an adjuvant.

26. A vaccine according to claim 24, wherein the immunostimulant induces a predominantly Type I response.

27. A vaccine according to claim 24, wherein the antigen-presenting cell is a dendritic cell.

28. A method for inhibiting the development of a cancer in a patient, comprising administering to a patient an effective amount of an antigen-presenting cell that expresses a polypeptide comprising at least an immunogenic portion of a colon tumor protein, or a variant thereof, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide sequence selected from the group consisting of:

- (a) sequences recited in SEQ ID NOs:1-1556;
 - (b) sequences that hybridize to a sequence recited in any one of SEQ ID NOs:1-1556 under moderately stringent conditions; and
 - (c) complements of sequences encoded by a polynucleotide recited in any one of SEQ ID NOs:1-1556;
- and thereby inhibiting the development of a cancer in the patient.

29. A method according to claim 28, wherein the antigen-presenting cell is a dendritic cell.

30. A method according to any one of claims 20, 21 and 28, wherein the cancer is colon cancer.

31. A method for removing tumor cells from a biological sample, comprising contacting a biological sample with T cells that specifically react with a colon tumor protein, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide sequence selected from the group consisting of:

- (i) polynucleotides recited in any one of SEQ ID NOs:1-1556; and

(ii) complements of the foregoing polynucleotides;

wherein the step of contacting is performed under conditions and for a time sufficient to permit the removal of cells expressing the antigen from the sample.

32. A method according to claim 31, wherein the biological sample is blood or a fraction thereof.

33. A method for inhibiting the development of a cancer in a patient, comprising administering to a patient a biological sample treated according to the method of claim 32.

34. A method for stimulating and/or expanding T cells specific for a colon tumor protein, comprising contacting T cells with at least one component selected from the group consisting of:

(a) polypeptides comprising at least an immunogenic portion of a colon tumor protein, or a variant thereof, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide sequence selected from the group consisting of:

(i) sequences recited in SEQ ID NOs:1-1556;

(ii) sequences that hybridize to a sequence recited in any one of SEQ ID NOs:1-1556 under moderately stringent conditions; and

(iii) complements of sequences of (i) or (ii);

(b) polynucleotides encoding a polypeptide of (a); and

(c) antigen presenting cells that express a polypeptide of (a);

under conditions and for a time sufficient to permit the stimulation and/or expansion of T cells.

35. An isolated T cell population, comprising T cells prepared according to the method of claim 34.

36. A method for inhibiting the development of a cancer in a patient, comprising administering to a patient an effective amount of a T cell population according to claim 35.

37. A method for inhibiting the development of a cancer in a patient, comprising the steps of:

(a) incubating CD4⁺ and/or CD8⁺ T cells isolated from a patient with at least one component selected from the group consisting of:

(i) polypeptides comprising at least an immunogenic portion of a colon tumor protein, or a variant thereof, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide sequence selected from the group consisting of:

(1) sequences recited in SEQ ID NOs:1-1556;
(2) sequences that hybridize to a sequence recited in any one of SEQ ID NOs:1-1556 under moderately stringent conditions; and

(3) complements of sequences of (1) or (2);
(ii) polynucleotides encoding a polypeptide of (i); and
(iii) antigen presenting cells that expresses a polypeptide of (i);

such that T cells proliferate; and

(b) administering to the patient an effective amount of the proliferated T cells, and thereby inhibiting the development of a cancer in the patient.

38. A method for inhibiting the development of a cancer in a patient, comprising the steps of:

(a) incubating CD4⁺ and/or CD8⁺ T cells isolated from a patient with at least one component selected from the group consisting of:

(i) polypeptides comprising at least an immunogenic portion of a colon tumor protein, or a variant thereof, wherein the tumor protein comprises an

amino acid sequence that is encoded by a polynucleotide sequence selected from the group consisting of:

- (1) sequences recited in SEQ ID NOs:1-1556;
 - (2) sequences that hybridize to a sequence recited in any one of SEQ ID NOs:1-1556 under moderately stringent conditions; and
 - (3) complements of sequences of (1) or (2);
- (ii) polynucleotides encoding a polypeptide of (i); and
 - (iii) antigen presenting cells that express a polypeptide of (i);
- such that T cells proliferate;
- (b) cloning at least one proliferated cell to provide cloned T cells;
- and
- (c) administering to the patient an effective amount of the cloned T cells, and thereby inhibiting the development of a cancer in the patient.

39. A method for determining the presence or absence of a cancer in a patient, comprising the steps of:

- (a) contacting a biological sample obtained from a patient with a binding agent that binds to a colon tumor protein, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide sequence recited in any one of SEQ ID NOs:1-1556 or a complement of any of the foregoing polynucleotide sequences;
- (b) detecting in the sample an amount of polypeptide that binds to the binding agent; and
- (c) comparing the amount of polypeptide to a predetermined cut-off value, and therefrom determining the presence or absence of a cancer in the patient.

40. A method according to claim 39, wherein the binding agent is an antibody.

41. A method according to claim 42, wherein the antibody is a monoclonal antibody.

42. A method according to claim 39, wherein the cancer is colon cancer.

43. A method for monitoring the progression of a cancer in a patient, comprising the steps of:

(a) contacting a biological sample obtained from a patient at a first point in time with a binding agent that binds to a colon tumor protein, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide sequence recited in any one of SEQ ID NOs:1-1556 or a complement of any of the foregoing polynucleotide sequences;

(b) detecting in the sample an amount of polypeptide that binds to the binding agent;

(c) repeating steps (a) and (b) using a biological sample obtained from the patient at a subsequent point in time; and

(d) comparing the amount of polypeptide detected in step (c) to the amount detected in step (b) and therefrom monitoring the progression of the cancer in the patient.

44. A method according to claim 43, wherein the binding agent is an antibody.

45. A method according to claim 44, wherein the antibody is a monoclonal antibody.

46. A method according to claim 43, wherein the cancer is a colon cancer.

47. A method for determining the presence or absence of a cancer in a patient, comprising the steps of:

(a) contacting a biological sample obtained from a patient with an oligonucleotide that hybridizes to a polynucleotide that encodes a colon tumor protein, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide sequence recited in any one of SEQ ID NO:1-1556 or a complement of any of the foregoing polynucleotide sequences;

(b) detecting in the sample an amount of a polynucleotide that hybridizes to the oligonucleotide; and

(c) comparing the amount of polynucleotide that hybridizes to the oligonucleotide to a predetermined cut-off value, and therefrom determining the presence or absence of a cancer in the patient.

48. A method according to claim 47, wherein the amount of polynucleotide that hybridizes to the oligonucleotide is determined using a polymerase chain reaction.

49. A method according to claim 47, wherein the amount of polynucleotide that hybridizes to the oligonucleotide is determined using a hybridization assay.

50. A method for monitoring the progression of a cancer in a patient, comprising the steps of:

(a) contacting a biological sample obtained from a patient with an oligonucleotide that hybridizes to a polynucleotide that encodes a colon tumor protein, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide sequence recited in any one of SEQ ID NO:1-1556 or a complement of any of the foregoing polynucleotide sequences;

(b) detecting in the sample an amount of a polynucleotide that hybridizes to the oligonucleotide;

(c) repeating steps (a) and (b) using a biological sample obtained from the patient at a subsequent point in time; and

(d) comparing the amount of polynucleotide detected in step (c) to the amount detected in step (b) and therefrom monitoring the progression of the cancer in the patient.

51. A method according to claim 50, wherein the amount of polynucleotide that hybridizes to the oligonucleotide is determined using a polymerase chain reaction.

52. A method according to claim 50, wherein the amount of polynucleotide that hybridizes to the oligonucleotide is determined using a hybridization assay.

53. A diagnostic kit, comprising:

- (a) one or more antibodies according to claim 10; and
- (b) a detection reagent comprising a reporter group.

54. A kit according to claim 53, wherein the antibodies are immobilized on a solid support.

55. A kit according to claim 53, wherein the detection reagent comprises an anti-immunoglobulin, protein G, protein A or lectin.

56. A kit according to claim 53, wherein the reporter group is selected from the group consisting of radioisotopes, fluorescent groups, luminescent groups, enzymes, biotin and dye particles.

57. An oligonucleotide comprising 10 to 40 contiguous nucleotides that hybridize under moderately stringent conditions to a polynucleotide that encodes a

colon tumor protein, wherein the tumor protein comprises an amino acid sequence that is encoded by a polynucleotide sequence recited in any one of SEQ ID NOs:1-1556 or a complement of any of the foregoing polynucleotides.

58. A oligonucleotide according to claim 57, wherein the oligonucleotide comprises 10-40 contiguous nucleotides recited in any one of SEQ ID NOs:1-1556.

59. A diagnostic kit, comprising:

- (a) an oligonucleotide according to claim 58; and
- (b) a diagnostic reagent for use in a polymerase chain reaction or hybridization assay.

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84

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 120agttccggcg tcttgggccc ctcttcaagt cctgcctgga gccgtggcc ctacacagat
 180cagagacgga gtatgtcatc cgctgcacca aacacacott caccaaacac atggtttttc
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 312

<210> 20<211> 420<212> DNA<213> Homo sapien
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120tcccatcgcc ttgatctctt gtatctgcga ccgccttttt ctctgcacaaa agcctgggat
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300ccactagac gngtgcacat tctccatana agtggncctg cttcacttca caccagatcc
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420

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240caaataccaa gaatttttgc gtatgtttat attgtatngt tctaaataat gggtagncicg
300tgaataaaga tcttgccacc catgtaataa tantagtaat actatagtna naaatggctg
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420gacaaagttat tttgt
435

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<210> 22<211> 407<212> DNA<213> Homo sapien
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60tgatacagta cgcacacgct cacttgaaag tctagaatc agaggataaa gaagccataa
120gccaccocac ttacatttcc tactatacaa tgcctttttg gcgcttgata aatcaagcat
180tcagttagcga ttacattcaa cagaaacatt tctcgtactc tgggtttaag atcctttgoc
240ctcagntcgc gatgtcgtga catctgactc tctcctcatt taaatatitt canccatttg
300cctatctcgc atgatgtttt cctcagacac tgagcaaatg acccaaggct nattgggggtt
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407

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<210> 23<211> 272<212> DNA<213> Homo sapien
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120caaaaacctg tttttgaatc cccaagaagg cagcatgtgt atacaacctat accacctgtg
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240attattacct gcccgcgccg cgtcgaaggg gg
272

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<210> 24<211> 424<212> DNA<213> Homo sapien
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60taaaccaaca taaaaaaggg aaaaacataa aagtctccct ttataaataa tattattcc
120agataaggaa gttgaacaaa aggtttgtta agtctctgca actatttcagt ttaataacca
180attatccctc aatattacaa aataaaatga ctggatcaat gntgactctt ctttgatctc
240attattaaaa tgctttgata ttacttacta agttccctga taactcaaac aaggttaaat
300taaacacttct attttcaggt attgctatct tagaagagga ttaccoactt catttagatg
360aaactgtttt accaanagcc tctcaggat ccaancana gnaattotta ttgcctctca
420tttt
424

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<210> 25<211> 372<212> DNA<213> Homo sapien
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120actcctcggg acccantgtg ccggagtgat ccgggtcaaa gtggttgaag gaggcccgga
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240cctcatgatg gncctggct gatggnggtg gagcatctgn tcccatccca ccggatgng
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360gtcgntcctc cc
372

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<210> 26<211> 342<212> DNA<213> Homo sapien
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60ggcctggagt ctattattca ccaaaagagc tgcacaacaa gtaagtgtc ctctcatgat
120atctgggaaa tgaatgaata agtactctgt ggctaccagt tgcattatgg agtcaactag
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240gaatgccctt gccagaagtn gaacatgagt aaaaattact ccaattgctt cttctaactc
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342

<210> 27<211> 315<212> DNA<213> Homo sapien
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120ggtttgggtt aacgtccgg gaattgcatt tgttttaaac cctaagtggg ggacagctca
180tgagtgcaag acgtcttggt atgtaattat tatacgaatt ggggctcaa tcgggagtcac
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315

<210> 28<211> 311<212> DNA<213> Homo sapien
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180gaggtggggc aggtacaact gtgtgagtga anctcgggct gtcaactggaa ggggtgnaat
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311

<210> 29<211> 516<212> DNA<213> Homo sapien
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120cacagagttt gtaatgattg ttgtggacag tacagacaga gagaggattt ctgtaactag
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480attaatggtt tagatatatt nataataaac tgattt
516

<210> 30<211> 355<212> DNA<213> Homo sapien
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120gctgcttcc tctgcgccag aagaaggccc acctgatgga gatccaggta aacggaggca
180ctgtggccga gaagctggac tgggcccggc agaggcttga gcagcaggtta cctgtgaacc
240aaagtgttgg cgaggatgag atgactgaac gtaatcggtt gaaccaaggc aaaggtcaca
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355

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240ctctctcat tgaactggtc attgaataat ttttttttct ttaagctatt gagacatggg
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355

<210> 32<211> 285<212> DNA<213> Homo sapien
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285

<210> 33<211> 250<212> DNA<213> Homo sapien
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 409

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 225

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 267

<210> 38<211> 556<212> DNA<213> Homo sapien
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 556

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 120gaggaggagaa gagattcgt tctgagctc ctactcccg gttctcgta gagaagccga
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 203

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<210> 40<211> 560<212> DNA<213> Homo sapien
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420gntcaccctc gtagaagntt ccattgtaac caaaagtcac aacccccacc gnttacacaa
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560

<210> 41<211> 265<212> DNA<213> Homo sapien
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265

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407

<210> 43<211> 343<212> DNA<213> Homo sapien
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343

<210> 44<211> 186<212> DNA<213> Homo sapien
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180cccttg
186

<210> 45<211> 503<212> DNA<213> Homo sapien
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60taataaacctt aaagggggaag tgccctgggt gactgtaag tcaagagaat ccactccagg
120aacagcactt ccaggccctt gcctttaaga actcctttaa ggcaagatng gacacatgcc
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503

<210> 46<211> 559<212> DNA<213> Homo sapien
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120cgagtgccogt gccataggcg gacaccaggg cagggttccc cgtttgtcca ngctccocaa
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559

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<210> 47<211> 513<212> DNA<213> Homo sapien
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300gtacctcact agagttaatc ccaccaatc agattgagaa ataaattggg gaatgtggaa
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420cctctacaat tattttcttt aagctgantg ctncctccct ggaatatcca tgttttctcc
480ctctcnggan tatntngcgm tccccccana cct
513

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<210> 48<211> 413<212> DNA<213> Homo sapien
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300atggggatct gcaacttoaa tcaatcaact tcatctggag ccagccocctc cggccocagg
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413

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<210> 49<211> 560<212> DNA<213> Homo sapien
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60tcaccacttg caatgagcgg ttcogctgcc ctgaggcact ctccagactt tocttctgg
120gcattggagtc ctgtggcctc caaaaaacta ccttcaactc catcatgaag tgtgaagtg
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420ccatgttcca ccgcaaatgc ttctaacggy attattgaat tanntggcgt tacacccott
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560

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<210> 50<211> 231<212> DNA<213> Homo sapien
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120agccaagggg ggtgtcattt totgggaatg gttaaacaca aaaggctgat agctggtatc
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231

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<210> 51<211> 265<212> DNA<213> Homo sapien
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120gttgaggcga ttctggggtc gctggatgct tacttgaa gaaggacttt tctggtgggc
180gaagagtgta cattggctga catcacagtt gttctgaccc tgttgtggct ctataagcag
240gttctagagc cttctttccg ccagg
265

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<210> 52<211> 318<212> DNA<213> Homo sapien
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9

60agaccgttaac agtacaatat ctttatgtgc acaatttact gcaattgtat tcagactcaa
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 180atgcccaaac aatctcatat gaagtcaccc tagccatcat tctactatca acattactaa
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 120gaagcctctt acaaaattat ttacatcgtt tgcacatctt ttacatcttt taagagcaac
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 240gttggtccccc ttggaaccaa tcacactggt tctcgtatgc tggggacaag taaaaagcat
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 360gtcgtgttga cttgtcaact aatttgaana aacgtcgttt ggctcactgg cgaagggtta
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10

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 120atttgactga agaatttttag aaaaccagga gagaagacct tcacccaacg aagcogctott
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 180tcgaagaact acttcataag actgttatga caatctgttt ttattacata taatttagta
 240gtcatgtcta atcaacaagg tcaacagaaag cttgtgaata ttctgctaaa taacttagca
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 180

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 120gttaacacgc ttttattgct attcaagtag caaaggaaaa ctactctcac aaacttcagt
 180tcaacagaga agaatcacca ttaagattga gatattgaa tgactaaaaa cgaagtctcc
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 420taaggacact ctgatgatcc ccacgaacta ngaggattgg cggtaggtcc ttatagatag
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 478

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 347

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 240tgacatttct ggtgtctctc ctgttctccc tcagcccctg cggccccacg gttgtctgtg
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 349

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484

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325

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255

<210> 74<211> 244<212> DNA<213> Homo sapien
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 570

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 180caattcctnta gtcaaaaagt ntcnctttg gaagcnctaa ttttgctgtt catgngcat
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 360taacnccnctn ttgagcagag ggaactanaa taatgtagcc atatacgggt nctcaaat
 420ntacgtatc tatctaaagg gggagcnaaa gggggcgga aaggacacaa ttttaacana
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 120agggagagc atgtacggtg tggggaagt gaaaaaaaag ctggcggggg agaaggaggg
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 120aatttgaaag gctaggagaa actcctccan cttaaaaaac tcaacttoga ccaggacagc
 180ctgacctcgg atcctggcct tctcatgacc tctgtgatc ctctctctct ctctcgtga
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 120ttgcctctc cctgataaga tttgtcccaa agggctcgtt aaggaatctg cccacaagt
 180tcccccatag aaggatttca tgagcagatc aggaacacta gcaaatgtaa aaataaaatc
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 318

<210> 87<211> 435<212> DNA<213> Homo sapien
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<210> 88<211> 293<212> DNA<213> Homo sapien
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 180ccaggacaaat cagtaaaaat ctacagtaac ctgatacaac aaaaatccct aggtgtgtg
 240aaattacaat ggtcacctct gttattctta aacttaaaat gatgtgtctca ttt
 293

<210> 89<211> 264<212> DNA<213> Homo sapien
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 120acctgaact aggaagaaga tgcttatctt gatgaataa tcaacagccc accacagta
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 120catgagcagc aactcggctt ctgcagcaaa cgggaaatga cagcaagaag ttcaaaggtg
 180acagccgaag tgcaggcgct cctctagag tgatccatc ccggaagctc cccatgcagc
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 120caacaaacag ggcaagtaat agcagcttcc tttttgtcac tctccatctc ccaagtggt
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<210> 93<211> 387<212> DNA<213> Homo sapien
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 180ggtattacat ctctctgga gctccattgc agttctggac ttaagaacc agtcaaatct
 240gaactgagga gagttggcaa tacactgtcg caattcatca taaacatttt ctttgtcaaa
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 387

<210> 94<211> 233<212> DNA<213> Homo sapien
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 120tatactagac ataacttccc cccacccagc ataattgtat gaaatattta gaattacaag
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 233

<210> 95<211> 268<212> DNA<213> Homo sapien
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 268

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 178

<210> 97<211> 338<212> DNA<213> Homo sapien
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 373

<210> 99<211> 344<212> DNA<213> Homo sapien
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 264

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 409

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 185

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 477

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 120cctgggggaa gtacaaacct tggcagtcac catcttccaa atagggtggg gtctggagct
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 575

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 410

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 146

<210> 108<211> 273<212> DNA<213> Homo sapien
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 154

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 344

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 469

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 415

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 420acgtctgtga cagttttata attgctctta ggtataatg gttagctaga tagtatactg
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<210> 121<211> 246<212> DNA<213> Homo sapien
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 120tgcaacttgc tcaagctgga atctccgag ccgccttttg cctttgcctt tccctgctgc
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<210> 122<211> 406<212> DNA<213> Homo sapien
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406

<210> 123<211> 596<212> DNA<213> Homo sapien
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<210> 124<211> 255<212> DNA<213> Homo sapien
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 120ttatttttctt ttgttttcca gaatacttat aattctttga gcctcccaga aattggaagc
 180taataaagc aactcaagtt tcttcaaaa aaaaaaaaa aaaaaaaaa aanttttnc
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 255

<210> 125<211> 332<212> DNA<213> Homo sapien
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 332

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<210> 127<211> 582<212> DNA<213> Homo sapien
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 120ggtgtggctg caccocagca ccgggggggg gcacgcancg acaccagagc aagggaacg
 180ccccccatg tcaacagcag gcaacctgtg ttttcat1tc aagtgggata cagtatt1tt
 240ttaataagga gcatacttt tttttaagag tt1gagatct gaatgtgatt tctaattgta
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317

<210> 129<211> 582<212> DNA<213> Homo sapien
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 420ctcatcttcc tgtagcngct gtgtgtgttg cntgaaact ctgtcncctg tctctcgggc
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 60gacgactctc tggggttggc agcagtagtt gcataaacat tgatgttato cggcag 116

<210> 131<211> 198<212> DNA<213> Homo sapien
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 198

<210> 132<211> 308<212> DNA<213> Homo sapien
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 308

<210> 133<211> 262<212> DNA<213> Homo sapien
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 262

<210> 134<211> 343<212> DNA<213> Homo sapien
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 180actgcactgt ctgagaattt ccaaaacttt aatgaactaa ctgacagctt catgaaactg
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 317

<210> 136<211> 159<212> DNA<213> Homo sapien

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159

<210> 137<211> 264<212> DNA<213> Homo sapien
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264

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263

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459

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576

<210> 141<211> 386<212> DNA<213> Homo sapien
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386

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227

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240acgtgg
246

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318

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295

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147

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401

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221

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551

<210> 158<211> 339<212> DNA<213> Homo sapien
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180cttgtgtgag cgtgtggaca agtgggtggc gncctgtgcc tgctcgtgtt gctcaatgt
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339

<210> 159<211> 385<212> DNA<213> Homo sapien
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120actcccgcca ggnatgncct acnctcanc tgtctttgga agctagtatg taagtaaggg
180aggagtcato aagtttatag atgggtaggc tgaggattga ggcaggaggg gacttaatgg
240ctagatccct gcttggttcc agagccctgg cccttgagcc cctggactgg tcaagtcatg
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385

<210> 160<211> 147<212> DNA<213> Homo sapien
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120taactgagt gtgnggcac gcnctct
147

<210> 161<211> 176<212> DNA<213> Homo sapien
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120gcatttgacc taaccaatgc gaaaaatent tncogtttna ttttttccaa atattc
176

<210> 162<211> 148<212> DNA<213> Homo sapien
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120tttgttttca gatccaactct gggcttct
148

<210> 163<211> 237<212> DNA<213> Homo sapien
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120ccgctgtaac tattctgnoc ngcattanto taagtentaa tgggcoctcc atccctacg
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237

<210> 164<211> 337<212> DNA<213> Homo sapien
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120acagtacaat ggaactntt naaactnttg nttcttgggg gcttntttcc aataacatca
180cttgatgcaa ctgggaacct ggtatttgcc aatgggggag gagcccccac catcgtgact
240gccctctctg tcttgaaccc tcagaaacct tctctgtcca ccagcaaccc tgttagcttg
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337

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120atacaaaactg ctcaactagt aagggaacaa aagaatataa tccatggtgt ctgctgattc
180aaaggggaga aacaaggntg tcatttaata tncnaaaacc
220

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<210> 166<211> 739<212> DNA<213> Homo sapien
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 120agctcctgcac ncaaggaatg gagggtctcnn acctcccnng ggggaaaaana ccongagtggtg
 180nctcncntcc ttgcctgggg gagnacaaaa anaaagncgt gctttcttgt ttttggaggg
 240cccaaacan agaatactgg gaatcttgga agcttccott ggtgtgaaag agaaacaagg
 300cganggttga aataattagn atataaatc actcaaggta atgctaagta atagggttga
 360ngtgaaccca gcaagctttc ctctganaaa gaaataccat caaanatcct tctaaaagct
 420tatctcctca gcatcattac attattttat ctctggtaat tcaatacgg agaagattcc
 480attccttgaa atgcactagc aaaatctgtta catctcagtg agttatccctg cccctttmct
 540ccaaaagtg ccaaagttta tgattacnng acataaaaata acaggttctg gaglccctgc
 600ttcaektga gaataaaggg taltgatagg ngctgnngga tggatgactc gtttctnanc
 660gtctattana nttggancgt ggggaaant ccttgcccta ggtcctgagt ggaaanatat
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 739

<210> 167<211> 290<212> DNA<213> Homo sapien
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 120ttttattaat ctatttaatn agggaaaaaa acnttnagaa tccataangt ttcatgttat
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 290

<210> 168<211> 250<212> DNA<213> Homo sapien
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 120gtgaagaaa aaaaacnaaga aggggaaggg aaccacaatt tatntttggg agtttttact
 180ggcactgtct caggacaagg ctactgttcc taaatacatc aagtggaacc agcgagagaa
 240aggcattttt
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<210> 169<211> 146<212> DNA<213> Homo sapien
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 120ttgctgaggg caacacnng ggtcac
 146

<210> 170<211> 292<212> DNA<213> Homo sapien
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 60tagataatt ttgaattatt tcaagtaagt agatgtttaa gacttgactc aagatgtgct
 120gaaaagaana ggaattctta ttccctgtat ctttttttcn cttnatgga onatatagtt
 180cttttttga agatgcactc attctgata ctottaaccg catttattct tactagcatt
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 292

<210> 171<211> 151<212> DNA<213> Homo sapien
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 120tttttggcgt aaaaanaacn tgtgtgtgnt g
 151

<210> 172<211> 131<212> DNA<213> Homo sapien
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 120gaagcgttca t
 131

<210> 173<211> 90<212> DNA<213> Homo sapien
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27

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 360gggttatatt gtctctcgac cactglatgc gggcctggg ttagctgttt gagtctcat
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 472

<210> 175<211> 752<212> DNA<213> Homo sapien
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 120gaggttgtag taagccgana acaccccctt gcnctccanc ctgggcaaca gagggagact
 180ccatctcaaa aaaacaacaa aacaaacaaa caaaaaaact ttgttttaa gtggtcgaga
 240ctatgtgcta gactttactac tgtttaatat gctaaaatga tacataattt attcttcaca
 300gtccaaaaatt caatgtaaa ccaaatatta attttgtcca gttagagaaat taaatgttta
 360caaacaaaagt atggaacagt tgttggccta tgtaaaatga aatgtaattg accattttat
 420tttcaaacat ctgtaactct cagctgtctt ttccactctt atactcattt tgcagaataa
 480agatgactgg tactaattta gacaataaac aaactacacc aagccacatt ttagtataag
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 720ntcgancntg catnttnang gccaatctcc ct
 752

<210> 176<211> 224<212> DNA<213> Homo sapien
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 120cgccgcccgg aaaaaangn gganaaanc ccgmagggt tgaagctggc ttcttcgaat
 180ttgcgaattc atatgaaat cactctggag ctggtaaaaa gagg
 224

<210> 177<211> 294<212> DNA<213> Homo sapien
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 120taactattgga tatggacctt ttgtntttgg gtgaaaactn caaagtaagg agacactgtc
 180aatcaattcc actaaaaatt catttatttt cctgtcatatc taaaaaaggg aaaaacagtag
 240caaatactggt gctctgtttt cccctcaac ggcaagcctt ccacaacagc acag
 294

<210> 178<211> 142<212> DNA<213> Homo sapien
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 60ttgcagcttt tgaagggtgg actctttcta ttgacacact ttacaaggag gattgtaaag
 120gtaattactca gtaccanaa ac
 142

<210> 179<211> 366<212> DNA<213> Homo sapien
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 120gggagttcta gaaccacttc ctgctncaag aagggggcct atgtctcgtc ggcttccagc
 180ttcaggacac agcatccacc ttgctctcgc cagtggatcc cctgcggtca ggctggcag
 240ccccanagag aggatgtgga aagcactttt tggctgacct catctggggt tggcaacagg
 300acaganttca caggaggcca gtggcggggc catgagggac agggtctcttn nncattcttt
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 366

<210> 180<211> 104<212> DNA<213> Homo sapien
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 60ntgctgaact ggggtntaac acctnctgg agactagact ccag

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28

<210> 181<211> 393<212> DNA<213> Homo sapien
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120tatgttgggt gagttcaaac acacnaaaaa cmcctntnt gncacaaact gtcttggctg
180gggttgggat aggtctgcat gctttttgaa gtttagtaca gctgtatat tcattaacgga
240attcagataa aatttcctta tgtttctgtg ttatgtttga tcgaatccta atcacaggga
300gtcttcatt agtcaaatat gcaatttgcc ctcaagtgcg cggctcatta ctttgaata
360tgccactgtg agtactgaca ttacagttg ttt
393

<210> 182<211> 311<212> DNA<213> Homo sapien
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120tgatactgc ctactctntt ctnaaaaaan gttagcnttt tcntnccagn gaaattattt
180ccataaagca aatgggtttct ctactctgaa aaccttgcta aaacccagtt ccagcataag
240ctgtctgccc acaactcaa tgtattgctt cattagagtt caattcatcc caatgagctt
300cacaggcaag g
311

<210> 183<211> 277<212> DNA<213> Homo sapien
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120cttttaactt ttctttttta agagtttctg ccaggtcttc aagcgttggg atatcttoga
180aaatttttat tcttttggcc aaaccaacat caccttcgaa cntttnttnc catcaagtca
240gcaatctgaa ttttgcata ctcttctccc attttta
277

<210> 184<211> 322<212> DNA<213> Homo sapien
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120gtgtacccat ctctgcccat cacccgttga atttngttg nctatttgga aaagatctgg
180ggactatctg aaactagtga gaatttacac caaacccaaa ggccagttac cagattaacac
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300caaaaaattct atcaagaatt tt
322

<210> 185<211> 358<212> DNA<213> Homo sapien
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60ttaagactaa gtgtacttga caattgaatg aattaagctt aaaaacattt ctctaagaaa
120ccagtggtcc atttaaccat ttgatgaaac nntattttta ttgacttata aaggatgtgc
180agtatactga aattccactt aaatactgaa atattctact aaatgacatt gttttgtcta
240aatttctccc agaaaaatct gttagcattt cttaaaagtc cctcagattt gagggaaatt
300ctaaaattagg acagttttct ctccaataa atataaatga tcttgagtat ttttgttt
358

<210> 186<211> 161<212> DNA<213> Homo sapien
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60cgggtcttc acgggttat tttataaaag aggaagaaaa aaaataaaag tctccggcg
120gggagacgcg gattttttgt aaattttttt ggggtttttt a
161

<210> 187<211> 408<212> DNA<213> Homo sapien
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120ggcaaggaaac tatatagaaa aacacatttg nttntntta aggccttactt ngggaataaaa
180ccattgttaca aattattgca catctgaaac cacagtgcat aacagactgt ctgcataaaa
240atgtctcaaga agtaaaacag gtatattacc tgacttagct cataaatgtt gatcggaaga
300caaatataga ttttcttgt caaagtatgc agcagtttga aaacttggcc ttcttgtttt
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408

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29

<210> 188<211> 195<212> DNA<213> Homo sapien
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 60tcacgtctaa tgagttaatg tctctcaact tctggtctca tgccatgtct cagtcaact
 120tcctgtagta cagcgattca aanaatntcn ttgttttncg ggaaacnnacc tgcctgggag
 180ggccgctoga aaggg
 195

<210> 189<211> 134<212> DNA<213> Homo sapien
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 120catggcataa ccaa
 134

<210> 190<211> 125<212> DNA<213> Homo sapien
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 120agttt
 125

<210> 191<211> 158<212> DNA<213> Homo sapien
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 120ggccatcaag ggtatgcata tncnaaaaa ncccaaaag
 158

<210> 192<211> 114<212> DNA<213> Homo sapien
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<210> 193<211> 147<212> DNA<213> Homo sapien
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 120acgaggggcaa cgaatccctcc ctcccaa
 147

<210> 194<211> 214<212> DNA<213> Homo sapien
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 120tatgtggatg tcttgatgag gaanctgggc ttgngtgaac aagtgcacca aaggaacgaa
 180gtgcgaatg aactgtaacc tgggcacatg tcag
 214

<210> 195<211> 296<212> DNA<213> Homo sapien
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 120cattgcagga gtgcaccggg aagtgcgaag agctgtttcc cattcagatg gaggggttca
 180agctcacagt caacaaaggg ttgagtnacc attttcaagn caaccacnca gtacgcctca
 240gcacacatgg ggagtccaac taccacttgc gggtcacata tgtggggaca aagcag
 296

<210> 196<211> 586<212> DNA<213> Homo sapien
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 120aaaaatgcct agcccacttc ttaccacaag gcacacctac accccttatc ccataactag
 180ttattatoga aaccatcagc ctactcaatc aaccaatagc cctggccgta cgcctaacgg
 240ctaacattac tgcaggccac ctactcaatgc acctaatgg aagcgccacc ctagcaatat
 300caacatttaa ccttccctct acacttatca tcttcaacat tctaattcta ctgactatcc
 360tagaaatgcg tgtgccttta atccaaagcc actttttcac acttctagta agctctaac
 420tgacgacaaa cacataatga cccaccaatc acatgcact catatagtaa aaccagccc
 480atgaccccta acaggggccc tctcagccct cctcaatgac tccggcctag ccatgtgatt
 540tcaactccac tccataaagg tctcctaact aggaactgac cgggag
 586

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<210> 197<211> 492<212> DNA<213> Homo sapien
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 60ggccgangta aacaatagta caaccctctg gtctctgttaa aactacatgg ttttaacacg
 120agctcaactcac aaaaatttttt tttttttaag taanacttcc ctgcaacacac agcannggag
 180gnaaacaaca ncaacaaaaa aateanante tgcagggggc ttgaaaaaac aggaagctnc
 240ncagtagtngg aaacccggag ctttttttta actttatatt ctttcccggt ttctcctttn
 300ntanaaacyn ggggtntctg ngnggccctc gtgtttggag ggaacggctg cagcggnga
 360anaaaactgc tgccctgggg gtgttggggn gggggtgtta tggattttct ctcccttng
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 480atccggggtg cc
 492

<210> 198<211> 414<212> DNA<213> Homo sapien
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 120acggagaagg acaggccatt gtaggagaag aggaacacca gctcggggat gtccaccacg
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 240ctgcctgttc actaagatgc tgtctcagc gctcaagttt gggggcaatt cttttgagag
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 360cagccgaatg ccaaggtctg cgcttgtttg actcccgtaa aagagagett ttgg
 414

<210> 199<211> 361<212> DNA<213> Homo sapien
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 120agacactccta gccaccaacg ggggtgatcca ctacattgat gagctaacta tcccagactc
 180agccaagaca ctatttgaat tggctgcaga gtctgatggg gtccacacgc attgaccttt
 240tcagacaagc cggcctcggc aatcatctct ctggaagtga cgggttgacc ctccgtgttc
 300ccctgaattc tgtattcaaa gatggaaccc ctccaattga tgccataca aggaatttgc
 360t
 361

<210> 200<211> 409<212> DNA<213> Homo sapien
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 60ggccgaagtc gccgccaa ca tgggtttcag gcgcttcgtg gaggttgccc ggggtgacct
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 409

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283

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634

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233

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240  atgtaagtaa  tgcattgctc  gctctcgaga  tcaaacactt  tcaggagagc  gctgtagtgc
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370

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240gtgttcagaat  caactgtgtc  atcaaaattt  aagtaaaatt  cagtgaaatt  aaggttgatc
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332

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 240

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 381

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35

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330

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300aaatttttagc aacagtatag aatagctcta tcgggtgggg agtaatcatt aaacagatga
360aatcggngcc agatttcat gtctctttag attcacagat ggaagcaaa c
412

<210> 236<211> 214<212> DNA<213> Homo sapien
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60gcgcgaagtct gagaagtgggt tcacttggta ctggatcatct actggcagtt ataactcaaa
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 180tgttaatca gaaagcatte ttncacttcc attt
 214

<210> 237<211> 176<212> DNA<213> Homo sapien
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 60caggtaaact cttctacatc cactaagtct taggaaaaac gtcaatcttc tgctgtttta
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 176

<210> 238<211> 526<212> DNA<213> Homo sapien
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 120ctgtcttccc cgcaaggac tatcatggc aaatgactta agctctctga gacaccatct
 180ccagattccc atccacttcc ccaaggattt cttgtctgtg ggtgcttgaa aaaaagaatt
 240tgtctgccat cgttttccct accgccatga acttggagca tccagagatg ctggagaaag
 300cgtcccgagg cgtgtggatg cgcgtcttgt caaggaatga agacatcacc gcgcgcgaga
 360cagctccggc gngtgcagag aaggtctgga tgctgcagaa acaagcccg gcacttcttg
 420aaaagatcgc aacgccaaag gtgagaacc agctcaagga gaccactgaa gcagctcgcn
 480gatnccggag ctttgggctg cccatcaccg tggcccatgt ggatgg
 526

<210> 239<211> 411<212> DNA<213> Homo sapien
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 120tgtctcattt agtcccatga aattaattat ttctctgtct tgatcttggt ggacagtttc
 180atgaagctgt cagttngttc attaaagttt tggaaattct caaacagtcg agngngtacc
 240agaaacttgt attcnagagt acaggtcaga gtcttctctt cttttctttt tgagattggag
 300tcttctctgt ttgcagact ggagtgcagt ggtgcgactt gggctcactg caatctccac
 360ctcccggtt caagcgatc tctgtcctca gcctcccgag tanctggggac t
 411

<210> 240<211> 319<212> DNA<213> Homo sapien
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 120cgacagggtt tteccatcat ctttcacggc gtaatgggca aagatgagcg tgaaggcaac
 180agcccatctt tcttcaaccc tgaaggggtt gccacaagtg acttctacc tgaagctgct
 240cctggccccc tccaccaaga agggcaaacg tcgcttgagc cctogaagtg tggcgctcat
 300ctccccgtac cggaaacag
 319

<210> 241<211> 97<212> DNA<213> Homo sapien
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97

<210> 242<211> 190<212> DNA<213> Homo sapien
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 60tgccngcgcn nccaagtngc ancgagaggg ngananttgc cctataccga ncncaagtac
 120aatgtcactg gcgcgcgcnt gacaacgttg ngaggccaca gcanccttgt cctccacggg
 180gttgaggtg
 190

<210> 243<211> 376<212> DNA<213> Homo sapien
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 60cggcagantt tggataacat ttgtattact acaagttgt tcttctgcg ttttctgaa
 120ccagtaaacg aaactcaaga ttgagctctc atgtaatgaa ttggggtaaa gaaaaaacat
 180cgagtgcaat aggttaggtt acaaaaggtt gttcacacat ttatgacag aggtcctnaa
 240ctgccaacac ctctaacat ctgattaggt tctatgagc caagtcttac atattccatt
 300catcatgacc ttttagtcaa tgtagcaaca cattttgcta aggaatggcc
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 376

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37

<210> 244<211> 405<212> DNA<213> Homo sapien
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 120caccatgaa ctccaccatc caataaactt aattggctgc ataactgggc gccaaaggcg
 180caacattaat gagatccgcc agatgtccgg ggccagatgc anaattgcc accagtagg
 240aaggctctctc tggtaggc an gttactatca ctggctctgc tgcagatt agtctggccc
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 405

<210> 245<211> 312<212> DNA<213> Homo sapien
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 60ccgggcaggt ccaagggtca tgatggcagg agtaatcaga ggtgtcttgg tgttgtgata
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 312

<210> 246<211> 634<212> DNA<213> Homo sapien
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 60ggcgaggtcc nacaaggga tctacgggtg ataaagataa gacggtgaga gagataaggc
 120tgatgtgttc aagccttgac agaccatgac agatagaaca ggaactgtgt cctgttaacc
 180atttatccct aacatctagc atagagtcca gttagtataa gccataaac ctttgagtct
 240cttggcaaga taagttaatta gcacagatta ttgtcactca ctgcaactca gcccttgagg
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 360acagatcatt gttggggtaa gaaatggaag ccagmgtag aaaaagtag aatgccatgg
 420ggttagaggt agcagaggtc gggacagaac ttgtctgttc tgcccccttt caccctcctc
 480gttctttgcc ttatgtccaa cccatcactt gctggggtag tcagcctagt tgaacaggtt
 540tagacaaccc tagagttctc tcacaggaga ttaatactga gaangagang tcttaccatt
 600gtcactctgg tgaacacagc attctnactc agag
 634

<210> 247<211> 325<212> DNA<213> Homo sapien
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 60cggcgaggtc cgggcaggta aaggcagaca ctgagtcagt attaatagat taactaaact
 120gcactgtaat ttagataaaa ttactgtgtc tcactgtgta ttacatgc aaatccacata
 180aatttgcatt taaccaacag tactgcacga gcgaactctc cgtatatga aaactgcact
 240taaatccaa cgttttggtta ctggaacctg catcataaat gcaacattgt catatgtgaa
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 325

<210> 248<211> 638<212> DNA<213> Homo sapien
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 120actaatgaag gctaagcaga atagtctgag ttgtctgaga ctaaagcagg gatagtggtt
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 240ggtgtcatcc aatgtgaagt gagcagtttc ggggtcaact ctatgtgtaa gaaactaaaa
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 360ctgatcatta aaacagagac accttcaact gtgtccata aggaattctct ttacaaaacc
 420gaacacaaac actcaaaaat ttaccatac ttgtaatgaa aataccagta ttgtgaagac
 480ncacagagc gggtttctat tagaacaagt atcagcaagg tcatgtagac ttgtgaagac
 540tttttgcctc tctctgcacca gacagatcat tgtccagcat tgccccggcg gccgctcaan
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<210> 249<211> 178<212> DNA<213> Homo sapien
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 178

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<210> 250<211> 477<212> DNA<213> Homo sapien
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240ccnagaatnc tggcgggatg ggttngnggc cgaagtggcag gagaggttga ggttcgctcc
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360cngggtgact gggctnctgc ggtttgcact cactgagttc tggnttccat atacatnngc
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477

<210> 251<211> 561<212> DNA<213> Homo sapien
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360acagtggcaa taatattgaa ctgggcacag catcgggaaa atactacaga gtgtgcacac
420ttggctctcat tgatccaggt gactctgaca ctattagaa catgccagaa cagactgggtg
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561

<210> 252<211> 284<212> DNA<213> Homo sapien
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120gagcangttg agtcttattt gttttatttt gctcatagtg actcttcagc agtgcaaaata
180ctctatctaa atccttcaag taattagtcc agtcaccacg actaagctctg tagttttgtc
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284

<210> 253<211> 656<212> DNA<213> Homo sapien
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180agaagggtca accacacac gtcatccatc ttgtctgaat ggtggggggc gttctcooga
240atctcgaata caatttggac ttctggagga aaaaacgtgca catgaaagac aagctctgc
300atctggcctt tgaggtgggc gcccgcaagg tgggtgtcct cctgtccaac ttgatcttcc
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480cacagttatg ntgcactctc caactgttca tccaccacaa cgtctttggg accnacttc
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600gcogtactag ttgatatgact tcgtcccaac ttggcgtatn tggcantaact gtttcc
656

<210> 254<211> 190<212> DNA<213> Homo sapien
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60ctccagcaac aactccaaac ccgtggagga caagtagct gtggccttaa cctgtgaacc
120tgagattcag aacacaacct acctgtggtg ggttaaatat cagagcctcc cgttcagtc
180caggctgcag
190

<210> 255<211> 446<212> DNA<213> Homo sapien
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60ctgagtcgta tgaatttccc tctgcacatt tttttcaga ttttcagtc gctttatgac
120ctacccaaca gaaaatgaga attaaaaaga atttgtcaaa ctatctttaa taatgccct
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300tatbanattt gcggaacgac gtggtggggg gtaaaaggca ccaaacctcg gccgcgacca
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39

420caaacctnggc gtaacatggg cataac
446

<210> 256<211> 315<212> DNA<213> Homo sapien
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60gatgtcttg tagtccagcc aacggatgtt ctogtcaaac ttitccatgt aggggttato
120cagcgcgtgat ttgtgagagt ggcctcccttc ttacgtccaca cctctgcagt gccgtgcgcc
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315

<210> 257<211> 524<212> DNA<213> Homo sapien
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120gaaactgtga acccgtaggg attaatgtcg gaaatggtta ggttttccag aaggggcagg
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240agctagacaa taaatcagat actagggggg agcctcaga tatgggcagt tegtccagca
300ccanagatat tataattccac agtctccagc aacctattg aatcaataat ttcaatggta
360aagtctctoga agatcccatc ngtatgccatc caggagagat tgaagctttt cgggagttat
420gtcagaaaaca tttaagtttc caatttcagc ttctttggct gtggaggact tgcctgggag
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524

<210> 258<211> 261<212> DNA<213> Homo sapien
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180tagaaaaaca ggaatagaat ggtgtgtttt atcatagtgt acacatttag cttgtgtgtaa
240atgactccaa aaactgattt t
261

<210> 259<211> 190<212> DNA<213> Homo sapien
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190

<210> 260<211> 692<212> DNA<213> Homo sapien
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120tgatgaattg ggaagagagc aaaccagaaa tggctttatt ttctccctgt gactaatttt
180taagtctcoga ttggaattca gtgagttagt tcataatgtg catgacagaa ataagcttta
240tagtggttta ctttcattta gctttggaag ttttctttgc cttagttttg gaagtaaaat
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420taactatata aggttttgag tttagctgaa aagtgtacag attataaat gatatattgt
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540gtgtgatgtaa cttagatgat tgtctacagc gttctcaagg gtttccaaag agtttttttc
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660gaactttggtn ccaactttgg cgtaatctgg gc
692

<210> 261<211> 356<212> DNA<213> Homo sapien
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60aattatacca aattctcttt atcaactgca tactaagtgt ttccaatata attttttcog
120tataaaaaa ctgggaaaaa aattgataaa taacaggtta gagaaagata tttctaggga
180attactcaga tctattggaa aaagttagta ctgtggatat ttaaatatato acagtataca
240gatcatcatt gtctctacag tatctggggc cagacactta agtgaagaga gaagtgtttg
300ggtgacttct ctacttaaaa ttttggctcat atcatttcaa aacattttga ctttgg
356

<210> 262<211> 366<212> DNA<213> Homo sapien
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 360tttgtt
 366

<210> 263<211> 389<212> DNA<213> Homo sapien
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 120gtgaagang toattattg agacagccct gttctccctg tcacctgca gtgtaacctc
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 389

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 300tgagggggc catggcaatg gcttggaact gccggcgcg ccgtctgag ggccgaattcc
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 409

<210> 265<211> 161<212> DNA<213> Homo sapien
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 120tgacacata atgaaggag ctggctgacc tcatgaccg c
 161

<210> 266<211> 455<212> DNA<213> Homo sapien
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 455

<210> 267<211> 261<212> DNA<213> Homo sapien
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 120gggactgtt tanagagcct gtcaccagag cttctctgg ctgaatgmat gtcattgtct
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 261

<210> 268<211> 111<212> DNA<213> Homo sapien
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<210> 269<211> 289<212> DNA<213> Homo sapien
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41

60cttctggaac agtcacottg ttaattttat ttttgaanaa tatttttcca ctctgcocct
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 289

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 60ctgggggtgg acagcgccat gtccctgac caggcgcaa agaacttgat gaatgcctg
 120gtgcagacag tgaaggcatc ctactgtgcc tctacocaa accaaagtc acagggtag
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<210> 271<211> 220<212> DNA<213> Homo sapien
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 220

<210> 272<211> 238<212> DNA<213> Homo sapien
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 120ctcagaaata tacatagaca aagttagcta atgaataaaa taagtaaaat gactacaata
 180actcaatttc agggatgagg gatcatgcat gatcagttaa gtcactctgc cacttttt
 238

<210> 273<211> 504<212> DNA<213> Homo sapien
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 240acaaacacca caattacact gatgcacct ctgaggggcc aagagacaa atgaagtgg
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 360acaggagaaat ctgcacaacc aatgaagggg tcatgtaccg cattggagat cantgggata
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<210> 274<211> 388<212> DNA<213> Homo sapien
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 120gacaaatgta aacctaaaga atgtgcagc gaatgcacaa agagttgtcc tgtagtctga
 180atgggaaat tatgcataa ggttacacc cagagcctaa tagcatggat ttccgaaact
 240ctttgtattg gttgtggtat ctgtattaa aatgcctcct ttggcgccct atcaatttgc
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 388

<210> 275<211> 344<212> DNA<213> Homo sapien
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 180gtgtatgagg gggaaatggt ggggtcgtct gggccataga ggacattcag gatgactgg
 240ctcgtgtggt caacacttaa ttgttctggt attccacact catagggctc taatacattc
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 344

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42

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 240ctgaaacaag aacacttggt ctacatcgaa cccctgcgag gtatltttca cccaagaaat
 300ttcggatttc aacaagagac ccattctctt ggataacaac gttgatgggg aagtgcagat
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 418

<210> 277<211> 758<212> DNA<213> Homo sapien
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 120aacctgtcat acattcatga taagtgcac tgaaaaatta ctcatioaaa ttccocctgg
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 420tttactgtga ctgatttga agcaaatata tactccaagt ccatgcagct gaatacacact
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 60aatancctac ttanttaatt gggagggaag tngtaaaaaa aacntttagt aatttgcgna
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 758

<210> 278<211> 392<212> DNA<213> Homo sapien
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 120tccacatggt gatttaataa agtcagaatg gcaggtggga ggatggtaaa ataactctac
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 360aaactatctc ttgcccacaa gaagcacatc aa
 392

<210> 279<211> 88<212> DNA<213> Homo sapien
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88

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 588

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 180ttggtactag gaagaagtct aggatgtgaa ttacatatat acttcccctca gtgactatga
 240taatacaagg ggccagatagc agaggaaaa atgtlaaacat gaaatttgac aaattttatt
 300actttgcocaa aattagocaa acaaaaatac tcaacctccc ctgtccaccc cccaactttt
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43

420aagatacctt atatgcocata aagttaatac cag
453

<210> 282<211> 708<212> DNA<213> Homo sapien
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708

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120gcctctctca aggatgaggt ttgaaagatt atgccagtgc agaagcacag ccgtgcccgg
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227

<210> 284<211> 478<212> DNA<213> Homo sapien
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360ggcccaactc cgggaattcca tcaacctcac caacctcaat ccaggccacag agtatgtggt
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478

<210> 285<211> 150<212> DNA<213> Homo sapien
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150

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328

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120atttctcgag cgtctgagat gttagtatta gttagttttg ttgtgagtgt ttagaaaagg
180cactacagga ctaggagga gataaggaaa atgattatga gggcgtgato at
232

<210> 288<211> 418<212> DNA<213> Homo sapien
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60ggtgacag cacatccggt atggccocct gaataccgat gatgccatct cctcagacca
120ggagagactt catctcctgt ttcccaagtg tgcatacaat atcagacot ctccagcat

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180cctcatgggc ttgctocaaq gggtttaact cctgcgccctg accagcctcg tgtgcgagaa
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 418

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 206

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 360

<210> 292<211> 174<212> DNA<213> Homo sapien
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 60gcttgccacc accagatgag aagttaagca gcctttctgt ggagagttag aataattgtg
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 174

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 406

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 120ttatctcagc atcaggtgac aagagaactg atatgggaga gattcagtga agatatggac
 180ttggaagactc caaggccgct tgtctttgag acctcagact cgaataagta tgcacaaagt
 240tggatgtcca ggtagcacc ctccctcag atgaccattg ctagcaagaa acaggaggcg
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 304

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<210> 295<211> 349<212> DNA<213> Homo sapien
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 240tgcattgttga gtggcgccctc tttggaattg caatttgsa aggatattga agaatacagt
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 208

<210> 297<211> 218<212> DNA<213> Homo sapien
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 218

<210> 298<211> 545<212> DNA<213> Homo sapien
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 545

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 410

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<210> 301<211> 393<212> DNA<213> Homo sapien
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393

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60attgatcttg tacaataata octgttaaaa aaatogtgct tacaacacgc toctanataa
120gagggcagggt ggagagagga cggagaaaac agctaccaa aaggggaggg ggagttt
177

<210> 303<211> 413<212> DNA<213> Homo sapien
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300ctgcacacta ctacaacatg gaagaacctc gaatatgtct agtgcaagaa gccagtcaaa
360aaagactaca tattatatga tactatttat atgaagtgtc cagaattggc aaatatatag
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120gtagaaaagg atgaaaaact tattgtctta atggaagaga tcatgagtga gaagagagaat
180aaaaacattg ttttttgga aacccaaaaga agatgtgat agottaccag aaagtattgag
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420gtattctga agatatcccc cangtttgc tgatctgtc gtgagggcag atcacattg
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47

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548

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318

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120ctaatacaat atgttgatc atggctataa taaagcagga gcaattataa aatcttcaat
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225

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48

180ggagagaata tgaactttact agcagagaaa tacaatatat cttgtotact ggaactgtaaa
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88

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100

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80

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86

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 244

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49

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344

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335

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295

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483

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50

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358

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306

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534

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282

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51

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 120ccatagttg ctaacaatcc tatttaacca ctaagaaagg atttacaaca aaaaaagcta

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52

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240ctgtgactga aacaataact cctgtctaa gntcagaagt aagatgcttc ttggggggct
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446

<210> 345<211> 197<212> DNA<213> Homo sapien
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197

<210> 346<211> 499<212> DNA<213> Homo sapien
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300ggataatatt ttctaatntt ttatttnaaa ttnttgctgc ttctttacct gcccgggggg
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499

<210> 347<211> 539<212> DNA<213> Homo sapien
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120gtttgtttct ttgtataggg tctgctctg tcatccagcg tanagtgcag tgaatgcaat
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69

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53

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251

<210> 355<211> 343<212> DNA<213> Homo sapien
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306

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357

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54

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250

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373

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276

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55

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540

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416

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173

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410

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541

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<210> 378<211> 368<212> DNA<213> Homo sapien
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 120tccagctcag tcccagcgct cagctgtcca atggcaacag gaccctcact ctattcaagt
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 120ctcaacgaag gcaaacacct ttacacgcta gatgttgggg acatcatcaa ogccctgttg
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 120gtcagtagcc tgaacagtgg cccaaaggcc actgatcaaa aataaaaatg tggctgtata
 180tcaatgaagt gaaatccaa agcgtttaac ctttggcat cagaaatcaa gattttccca
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58

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227

<210> 384<211> 218<212> DNA<213> Homo sapien
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<210> 385<211> 100<212> DNA<213> Homo sapien
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100

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207

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508

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120tgacattctt gaccatttgt gtgtttgttg ttgtactagt ttttgtttt tttaatgtag
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240tcagggtctt gcaactggac tgcagacct accaatcagc ggcattttat ctctctgaa
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351

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59

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 264

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 420gtctgtggtt atcccatgtt ggaattcatc ctttgtaacc cattgtccta tatcctaaca
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 240gtatgttata caagtcaaac ttggaaggtc atagtgaaga tactatgct gagagaaaag
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 366

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 544

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 120aaggtgtgca gcaccagggc agcccaaaagc agggctcggt tgaaaatacc aaagatctc
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60

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 432

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 120agatgtggtc atcaccatcg ccaacaatga tgcgatctg atcaccagga tctcagaccg
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 282

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 316

<210> 401<211> 469<212> DNA<213> Homo sapien
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61

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512

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130

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326

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224

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255

<210> 408<211> 643<212> DNA<213> Homo sapien
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643

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62

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 239

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<210> 415<211> 342<212> DNA<213> Homo sapien
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 180ggnaggtgta ctacgtgcct atgggggta tgatggtag gcagtcttt gccattggag
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63

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342

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286

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180ctcttcttca accatccat gttagctcta aggaggtttg aataagccat ctgaacttta

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64

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<210> 428<211> 365<212> DNA<213> Homo sapien

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 539

<210> 433<211> 539<212> DNA<213> Homo sapien
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76

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527

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 120ttatcttcta gttcaatggc gtccaggaca ggaagcctg ccaggaaacac aagcaggccc
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 420gaaactggga aggggcaccc agtgttgggc agaaaggaaa gctgggggttg
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 532

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68

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222

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376

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250

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120aaagacgcta atcttccaca ttgaaatcaa tgactaaaca tttttgattt acccagctac
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413

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328

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 359

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<210> 479<211> 562<212> DNA<213> Homo sapien
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73

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 360ccaccgaact ctggcaggat cgcangacg gatgggaatc cgaaaacctg gagatgaacc
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 180attctctcat accccacact aagttctgtc aaatctaccc taaaagtata tccactgttc
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 124

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 420aattttaagc atcaaatatt ggcncogaat ttgggaaaaa ttnaagnatg aatccttggg
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 564

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 327

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406

<210> 486<211> 386<212> DNA<213> Homo sapien
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386

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213

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297

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347

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480ttctaatgtct ggnatttctt ttttaccaaa ttgangaaaa tggaggnttg
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566

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561

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380

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535

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523

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120cttcaacggy ggagacctgg ttgggtgtct cacagagat gttttccggc tttaggtccc
180tggggcgat gcctttgtta tgcagaaagt ccaaggcgct gg
222

<210> 497<211> 86<212> DNA<213> Homo sapien
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86

<210> 498<211> 310<212> DNA<213> Homo sapien
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 334

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77

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397

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 496

<210> 513<211> 630<212> DNA<213> Homo sapien
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 214

<210> 515<211> 196<212> DNA<213> Homo sapien
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<210> 517<211> 338<212> DNA<213> Homo sapien

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79

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 180gcgcaaaagg gatgaagcag ttagttaact tttgtctga acagtccaaa ggaatatggt
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 319

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 120aggggtccag cctcgttga agaggaacag cactggggag tctttgtgga tctcaggcc
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<210> 521<211> 509<212> DNA<213> Homo sapien
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 343

<210> 523<211> 369<212> DNA<213> Homo sapien
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 120cagataggt ttggagtccc tttttcgata ctacagttat ggctcgaaa agaagtccg
 180cgtggacata tcaaggattt ttcaggagga aacggtgaag gactatgaag ctggtgaag

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80

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 353

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 180gtgagtgtn gctgtgttta ntgtgtgtgt gtntgtgnag agcaggagtg actgggnnct
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 272

<210> 526<211> 653<212> DNA<213> Homo sapien
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 120tgcccaactac ggaaaagcca acagtgactg tgaactccg aaagctgttg ttgaatcgat
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 240agatggatga agctgctcgg cagangaacn angacgcgtg aaggaaagct ggaanagctt
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 223

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<210> 529<211> 357<212> DNA<213> Homo sapien
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 120ttctatgaa tgttatcaat gtggaaaagc cttctgcoga agtctatccc ttattogaca
 180tcagatcatt cacacaggag agaaaacctaa taaatgcagt gaatgtggga gatcttcaa
 240ccgaagtaca aaacttaata agcatcaaaa acttcatgct gaagcaaaag acctgcoccc
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 357

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81

<210> 530<211> 179<212> DNA<213> Homo sapien
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 179

<210> 531<211> 288<212> DNA<213> Homo sapien
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 120gtctcatataa cttatgaaca gaaagtgtg aaataaagg gtactcatgg aaacacgtga
 180agagaggaaa caccggcaat tgttcaaac ggaacagtga gcaagtaact tgggagtaag
 240gtctcgagag atggaagacg ctggtctcag atctgaggng atgtctgg
 288

<210> 532<211> 320<212> DNA<213> Homo sapien
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 320

<210> 533<211> 578<212> DNA<213> Homo sapien
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 578

<210> 534<211> 457<212> DNA<213> Homo sapien
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 457

<210> 535<211> 394<212> DNA<213> Homo sapien
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 394

<210> 536<211> 324<212> DNA<213> Homo sapien
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82

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324

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314

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160

<210> 539<211> 401<212> DNA<213> Homo sapien
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401

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328

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615

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448

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83

<210> 543<211> 170<212> DNA<213> Homo sapien
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170

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572

<210> 545<211> 70<212> DNA<213> Homo sapien
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70

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427

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359

<210> 548<211> 362<212> DNA<213> Homo sapien
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360tg
362

<210> 549<211> 318<212> DNA<213> Homo sapien
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318

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555

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490

<210> 552<211> 197<212> DNA<213> Homo sapien
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197

<210> 553<211> 484<212> DNA<213> Homo sapien
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484

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200

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324

<210> 556<211> 349<212> DNA<213> Homo sapien

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 180tactcagttg tgtgtggttg taagggaact ggtgaagagt ggggttctgg gagcogatgg
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 314

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 321

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<210> 561<211> 330<212> DNA<213> Homo sapien
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 348

<210> 563<211> 325<212> DNA<213> Homo sapien
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 325

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 172

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 510

<210> 567<211> 319<212> DNA<213> Homo sapien
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 340

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 330

<210> 570<211> 371<212> DNA<213> Homo sapien
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 371

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 314

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 272

<210> 577<211> 509<212> DNA<213> Homo sapien
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 250

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115

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89

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432

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420tttttaatat ggatatgcta taaaactnct gnncccttta taattccctt ggccttgctt
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568

<210> 586<211> 345<212> DNA<213> Homo sapien
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180gggggtgaaaa gctcatccc ttanggttag gcagaaagtg taattgactt ggtatacacc
240actcttattg ggtgtgtgtc tgggcttata anctaactta ttttctcgcg accctagcat
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345

<210> 587<211> 116<212> DNA<213> Homo sapien
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116

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360

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461

<210> 590<211> 492<212> DNA<213> Homo sapien
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90

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492

<210> 591<211> 377<212> DNA<213> Homo sapien
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377

<210> 592<211> 401<212> DNA<213> Homo sapien
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300gtacagcgc atctcttcaa tcagtncagg gnccgngctn cagctcccan cangatgcnt
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401

<210> 593<211> 377<212> DNA<213> Homo sapien
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180ttaatgtctt acgtctcgc atttatatta aaaaattaca cacaaatgaa aatggaaaaa
240ctggccaatc ctgattttcg tccctattt ttcacatgc aatcatatcc tttagtcaat
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377

<210> 594<211> 310<212> DNA<213> Homo sapien
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180ccagaagtgt gactggctaa agctcgatgt ggtcacact gtatagctgc ttccagtga
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310

<210> 595<211> 434<212> DNA<213> Homo sapien
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240ctcttgagccc catgcgtttt cgatggtctg gctctggctc agatgccag ccgangaagg
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434

<210> 596<211> 740<212> DNA<213> Homo sapien
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740

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91

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<210> 598<211> 363<212> DNA<213> Homo sapien
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 363

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 488

<210> 600<211> 259<212> DNA<213> Homo sapien
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 259

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 386

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 120gtcgtacc tgaacagtgg cccaaggcc actgatcaaa aataaaatag tggctgata

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92

180tcaatgaaat gaaatccaag aagotttaac cotttggcat cagaaatcca gatttttcaa
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 317

<210> 603<211> 378<212> DNA<213> Homo sapien
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 240cttgaggagc tccangaact aggtntctcn ottttgaann gatgnaoat gcctnagatg
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 378

<210> 604<211> 359<212> DNA<213> Homo sapien
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 222

<210> 606<211> 507<212> DNA<213> Homo sapien
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 326

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 336

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<210> 616<211> 381<212> DNA<213> Homo sapien
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381

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315

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182

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133

<210> 620<211> 178<212> DNA<213> Homo sapien
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178

<210> 621<211> 280<212> DNA<213> Homo sapien
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280

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311

<210> 623<211> 269<212> DNA<213> Homo sapien
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269

<210> 624<211> 365<212> DNA<213> Homo sapien

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95

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 365

<210> 625<211> 391<212> DNA<213> Homo sapien
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 424

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96

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 600ctcctctttt gaatgggaac naagaaagtg aanaetono gatgntctcn actgcngga
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 720at
 722

<210> 633<211> 438<212> DNA<213> Homo sapien
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 120gtttagcgcat gaggacotaa aaaaagctgg attgctgatt ttgtctaata aacaagatgt
 180taaaagaatgc atgactgtag cagaatatcc ccagtttttg aagctaactt ctattaaaga
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 258

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 359

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97

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<210> 639<211> 261<212> DNA<213> Homo sapien
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 261

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 180tagcacacaa cagaaggacc octcccagc amnacaaca tccacttcca attaccagt
 240gcactnctgc tccctaaang aananacac acanacaac acnacaacac acacacacac
 300tca
 303

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 180aatgtgccta ccagattctg gtacaggtca aagaggtctc ctccaagctg agcacgctcg
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 295

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607

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223

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402

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109

<210> 647<211> 177<212> DNA<213> Homo sapien
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177

<210> 648<211> 240<212> DNA<213> Homo sapien
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120ccacgtctgc aaaaactccc acgtgctcta agaagagatg cagttccgg tgcctcgac
180gttgccagct ggctcgaac agtgccagga agatgtctc cagcatctcc tggaagtgg
240

<210> 649<211> 501<212> DNA<213> Homo sapien
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99

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 412

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100

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327

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824

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124

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135

<210> 660<211> 589<212> DNA<213> Homo sapien
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120taactgaact gatcaaggaa gtgctgctct ggaggagaag atccaogact tggccctgga
180gtcttcaacg cagagatcgc tgctggtgat gggggggggc tacaactatg ccaactgoot
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589

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120caggggcagc aggataagga atagagtggg ggcagaaagg tgggttatta aaaaagcatc
180ttgttactt gacatgaag ccacgtgccc cagcgaagaa gacagatga aggaactcgg

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101

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251

<210> 662<211> 654<212> DNA<213> Homo sapien
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654

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180gtggccctg gttcacacca taaaggcgca caagtgttca gtccatgaaga gtcggaagct
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330

<210> 665<211> 171<212> DNA<213> Homo sapien
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171

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636

<210> 667<211> 742<212> DNA<213> Homo sapien
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180aactccagaga ctgctacaag gaggaagaac acctcaactc gtcaatctgt ccaaccccta
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 543

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114

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 177

<210> 673<211> 439<212> DNA<213> Homo sapien
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240ttgtataaaa aggagagaag gatattcctg gactgactga tactacagtg octgcgcgcc
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439

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168

<210> 675<211> 406<212> DNA<213> Homo sapien
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406

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222

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530

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582

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104

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434

<210> 680<211> 412<212> DNA<213> Homo sapien
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412

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192

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458

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279

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497

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105

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 454

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 300tataaaatag tataatgatg ctgatgtcgt taaccaaagg gcagaataaa taagcaaaaat
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<210> 691<211> 102<212> DNA<213> Homo sapien
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102

<210> 692<211> 407<212> DNA<213> Homo sapien

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106

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446

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120actagtgaga cccacatttt gccacatccc acttcatggt gacaggagcc ctggtctctg
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263

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594

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402

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162

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120catggcccga cagggttaac tgcctctct ttgtatgtat ctggaagcca tagtttctct
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240ctgectctc agccactgaa tccctgtagt atctcaggct ttgatggcg aggacaaacc
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 360aacaggctgg cgttnacggg gntcctgaet ctgtcangat tggntttngt gtgaacaggg
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 526

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 120gcctctctgt ncagccaccg ctggactttt catnaaatgn cacagttctt ggtgtnttc
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 549

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 120aacctgttct caaggtagg gtgggaagtgt gtgtgagtgc gtggggggga gagggttgga
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 238

<210> 701<211> 500<212> DNA<213> Homo sapien
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 120ggctgaagg gaaagaagga ctgcttttgg gaggccctct tcactgcag ctatgatgoc
 180cttcccttct tccctgttcc tcacataatg ctttatccc attctactcc cctgtatgoc
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 500

<210> 702<211> 452<212> DNA<213> Homo sapien
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 120tgtcttatgt aaataataaa ctaattgtgg ctgtgaaatg atttgtatgt gatcctgtct
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 240tgcgggaata acaccaaatg gaatctctc atctctgtct tgttagogat gtgtctgatt
 300caggccatct gtctttttgt tacttttttg tccgtgtctc cctattggc ttttgtaact
 360gcaattttca aaccaaagta ctggcggncc ctagggaat caccctgggc gtctatgate
 420antgcacact gggaatctgg ctgtgtttct gg
 452

<210> 703<211> 286<212> DNA<213> Homo sapien
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 120cttacctgat ccgggtgpc acggaggaac caggagcgtg ccttcggcac cgtctggagc
 180ctcagagatg acaaggagca gctgtggaag aacacatat tctgttgacc gccctgtgc
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 286

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 60ttccctctgc cagataaaa actctaacca catacgggtg gtcagcttct tctttccca
 120actcagagaa agtcaataaa agggaaaaaa gaaaacatgc ttgaaaacac agtgaccaaa

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180ggattttgaag taataattac attaataaac cataactttt catttaacta ttcacattcc
 240acacagtggga aattatcctc tctccagat ttttcaacta caactctaac tttgaagacc
 300tacagttaaca aaaaacaact tacagacttc caggatgtgt gttttttttt ttttaatgcca
 360agcaacaagg gtcatcataa anattttgtg ttgatattta ttgatgaagc accaattttt
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 459

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 120cagccgctcta tgtggcaagt tgaagagagg aaggcagggg ggtcccgcca tgtcagccta
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 120aaaaaataaa acanctgtga tgattgtgan caaaatggca agtaagttaa gcatttttga
 180ctctgnaatc ntggmatcat tncaatgaaa ggaattccaca aactactcgg aaaggaagtt
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 360tcaattngaa tttngtact gatctggnac ctttactctt cttggagtta ctaattatgaa
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 472

<210> 709<211> 411<212> DNA<213> Homo sapien
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 120gaaagatcct catgaattaa atagttgatg caatttttaa cgtttaattg tataaaaaaa
 180aaaaacaaaa aattaggctt gtaaaaactga ctttttcaat acnngggttt tgaaintari
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411

<210> 710<211> 418<212> DNA<213> Homo sapien
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418

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526

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360g
362

<210> 713<211> 307<212> DNA<213> Homo sapien
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120taaaagcttc tcntacaga agaacagaca tgacccaac ttgttagtat ggacagttgt
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307

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110

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433

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341

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445

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453

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240tgagatgctga ngaagtgcgt gtagacatgaa gagacotgcc cgggcggccg ctcocgaagg
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 120ataattacga ttaccatttt tcttccataa tatatagcaa aaattctt
 168

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<210> 728<211> 564<212> DNA<213> Homo sapien
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564

<210> 729<211> 253<212> DNA<213> Homo sapien
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253

<210> 730<211> 291<212> DNA<213> Homo sapien
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180tcagatcatt cacacaggag agaaaaccta taaatgcagt gaatgtggga gattcttcaa
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291

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197

<210> 732<211> 203<212> DNA<213> Homo sapien
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512

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180

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 302t
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 463

<210> 737<211> 344<212> DNA<213> Homo sapien
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 313

<210> 741<211> 589<212> DNA<213> Homo sapien
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 282

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 271

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<210> 754<211> 466<212> DNA<213> Homo sapien
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 120ctcctgcctt ccatgatgac agctgctgtca agtctgtcgg cagggtcaca gtctcttaga
 180gtagacttca tcaccagctt ggagtacttg tactcatgt ttactctct ttccaaaag
 240tagcagggtca agaccgtgag cagggtggca gtacagggtc ctgcatagat gccacatttc

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116

300agccagaat ctatggtttt gcanaatgga gactctctgc tcangonaga taaatncoan
360ccaaagcatt agcnttgggn ttctcnccnc caagttaagt aacnctttc ttgggaatcc
420cnmnacocca ccaaganttg gnttgaacga aatacctant ggtgta
466

<210> 755<211> 469<212> DNA<213> Homo sapien
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120actccctttt ctaaaactga actgaccac atcaaaagtt tgaatacaaa tctcatggtt
180aattaaactt gcaatcaaca ccatatggta acagaagatg gcaagagata agattcagat
240cttagatatt tccaagttag gcatgttaga tgatagaagg attagttaga agctgtagct
300agagctcagg ttgggcatga aggaactagt ctcccatgtg gtttggaaga natagggctt
360ccctganctc tattgtgaac tatacngggt tcatcccaag gaatggatga ngtgggcata
420aaaccttctt caaacctgaa ganggncccc ttntgtacca gaacacta
469

<210> 756<211> 412<212> DNA<213> Homo sapien
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120ggtacgcca gccccggaga accncgatgc tgacttttnc caggatctcc tctgggaatcc
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240gatctccccc tcactgtaga aggccccta caagcccaan atgtacngcn gacttgcat
300nngtgcaaga cctgcaagat cngnggatgat ntggttcnng atggcngctt tgactcgaag
360gtgcgataan attctngac cttgnggcgc gacnacnntn aagggngaatt tt
412

<210> 757<211> 385<212> DNA<213> Homo sapien
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120aaagtcttcc ccacggtccc ccaaatgggg gactatgggt tactgtgatc aagagacacc
180tgcaactaaa acacaaactc acttctacca aatcaaaact caaatccaca caacaaaaca
240gaattgagca atcttaccag ggattgaaaa ctgaggggtg gagatgctgg gctgagggcc
300aaggaggaga gaaggaggag agggagggaa agcacaataat gggaggagat gagggctcca
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385

<210> 758<211> 290<212> DNA<213> Homo sapien
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60gggaatgact tgagtttaga tacagatat tgcacacaaa aaagtcatcc tacaatgaat
120ctctatgaat gttatcaatg ttggaaaagcc ttctgcocga agttcatccc ttattcgaca
180tcagatcatt cacacaggag agaaaccta taaatgcagt gaatgtggga gattctccaa
240ccgacgtaca aaccttacta agcatcaaaa acttcatgct gaagcaagg
290

<210> 759<211> 288<212> DNA<213> Homo sapien
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60ctgttaagaa tgttataat gcaaatgatt ggaagcaggg catttttacc aaagagcaat
120gtcaactataa cttatgaaca gaaagtgtgt aaataaagg gtactcatgg aaaccagtga
180agagaggaaa cacoggcaat tgttcaacac ggaacagtga gcaggtacct tgggagtaag
240gctctgagag atggaagacg ctggtctcag aactgaggtg atgtctgg
288

<210> 760<211> 432<212> DNA<213> Homo sapien
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120gaaggagatc ttacaaagag taagggaaaag ggaagggggc agaggctgct tctcagagcc
180accagaagac aaaaataagc aggtgtgagc ccagtggagg aggcacgggg cagagacagg
240caactgttgc tggcagctg gtgcacgtag cactgtggga gatgaacctg gagaggaact
300agagaggaca gcacaaatgga gccaaagaa gacttagcat ggcggggcgc ggtgttcat
360gctgttaato ccagcatcttt gggaggccaa ggtgggcaga tcaactgagg tcaaggatctt
420gagaccacct gg
432

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<210> 761<211> 246<212> DNA<213> Homo sapien
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 60aatagggaac agatccctggg agccagagtc tcccccaggt accccaaagg ggacaggaaat
 120cggaaicgtg aagcgggaag ggtcttacct gctgggtgtc tggggccaag agactcgggga
 180agcacagatt ctgcttctca ccccaaacgg tgggggtggg ggtgggctga gatgcagacc
 240ctctgg
 246

<210> 762<211> 411<212> DNA<213> Homo sapien
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 60agctccttag ctggctgggg tggggagggg gtatgtcacg tggcagctgc tactcactgc
 120tcagtgtgga aaacacagga cttggcaatc acagcccga gaacctcat gtgtggcaga
 180agcctgaggg atcggttttc ttgcccagct gctctgttca ttttctgttg ttttctgtca
 240cttaagaat tcacatggaa gcatgtttta taaatgaat taccagagaa acagagatgg
 300gccagatatt tcagaaatgg ncccatgtga ccaagtctct ctgtttgggt gacagtgtct
 360tgaanattct ctttgangat gtgcantctt ttttttttt tnaaaaanaa a
 411

<210> 763<211> 581<212> DNA<213> Homo sapien
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 60atttttcaat ttttattttg gtttttttac aaaggttgac attttccata acaggtgtaa
 120agtgtgtgaa aaaaaaattc aaatttttgg gggagcgagg gaaggagtta atgaactgt
 180attgtcacaa gctctgatca atcctctctt ttctcttttg ccacaaattt aagcaagttag
 240atgtgcagaa gaattggaga gatctagctt tcagttaaaa aagaagaaga aagaattgca
 300aagagaagaat ttttcaaat ttctttcttt ttttaattag attgagtcca tttatttgaa
 360acagactggg caaatgtcca caaagaattc ctggtcagca ccacccgatg tccaagggt
 420caattacaan gaaaggcgag gctgtatggc ttattgtttt tgtattcaat gatgcttct
 480ccattcaatt tgtcttttta gagcagccat ctacaanaac agtgtaaagt gaacctgctg
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 581

<210> 764<211> 253<212> DNA<213> Homo sapien
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 120cctgagtcac acgaactcac cagagtcacg ggcccagact gggcctggg ctatggcgcc
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 240agggcgccct tgg
 253

<210> 765<211> 270<212> DNA<213> Homo sapien
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 120cgatggcctt cattacttac gtgctcctgg ctgggatggc actgggcatt cagaaaaagg
 180tctcccgcga ggtgctgggc ctgtgtgcaa gcacagcgct ggtgtgggtg gtgatggagg
 240tctggcctt gctcctgggc ctctactcgt
 270

<210> 766<211> 449<212> DNA<213> Homo sapien
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 60agtgtctggg aaaagtcaag atgctgacgc ctaatggctg ttctagcctt tccaggtttg
 120taacatgaag atggggaagg aaatggcacc actgctgttt gtaactcgag gaactcttgg
 180acgattccac tctccaaagc agtacaaaac ttacaaagaa gtcaaaagtc ttaacactcc
 240caattctccag gaactcttgt ctgtgtccat tggtagagg gaggaatctt ggttccctca
 300gtgccttctg atgttanctt ttgatagct tcaatccac tcnctgcct caactctgt
 360tctggcctg aatgtaataa ntgggtgtgc natcctaan naatcaactt tgnaaaagggt
 420tccctggac attcctcttt aacccnct
 449

<210> 767<211> 466<212> DNA<213> Homo sapien
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 60tcaagacca acaggagcat gtggtagcca cgtcacaaac caagaccatg gggcatagg

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118

120acaagaaga tctatgtggg cagtggtccc cgttaggctg cctcatccgg atattgattg
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 240aggcggtgat gccgataaca agaatcgccct ttaagggtcca agcagaagcg cctcgcgaag
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 360gatnaccgaca ctgtgctcaa ncaatnttag ggcctctatg aataaacttg acaataaann
 420anggggaacct gntaccoccca nncaaacccc gagcaatggg cttnngg
 466

<210> 768<211> 459<212> DNA<213> Homo sapien
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 120aggcggtccag cctcgttgga agaggaacag cactggggag tcttttgga ttctgagccc
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 240agatcctcggg tactgaaagc cttagggaag ctggcctcag aggggaagcg gccctaaggg
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 360catgagggaag gaacagcaat ggtgtcagta tccaggcttt gtacagagtg cttttctgtt
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 459

<210> 769<211> 409<212> DNA<213> Homo sapien
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 120actggggagaa tattgtaggg gataaagtcc ttgagaacaa acatcacact gaagcactc
 180tttncctggat cttgaagacn cacctctcca cctctaantg ttncctgttna tccatnaact
 240aangggccctt tttnaagaga tgtcctcctc ccttagtntt actctaacna cttctatat
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 409

<210> 770<211> 427<212> DNA<213> Homo sapien
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 120ctgtgattggt ggttcattat gtcttgctgt gcaggtagaa ggcttactag aagtgtgaaa
 180acntaaggctt ggattaaagg gacacccatt tctagatag tcagtagaat tagaattgtg
 240aagatgataa gtgtanaagg aaggttaatg gttgatattg ctagggtggc gcttccaaat
 300aaggtgcata gtaggtggcc tgcagtaatg ttanccggtt agcgcacgg cccagggcta
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 420taggnngg
 427

<210> 771<211> 524<212> DNA<213> Homo sapien
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 60cctgaccacg aactggggcg gctctcacto catgaggtat ttcttcacat ccgtgtcccg
 120gccccggcgcg ggggagcccc gcttcaatgc agtggcgctac gtggacgaca cgcagtctgt
 180cggtgtccag agcagcccgcc cgagccanag gatggagccc cggggcgccg tggatagagc
 240angagggtccc ggagtattgg gacggggaga caccgaaaagt gaagccccc tccagagcto
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 360ccgtgccaaan gatgtatgcc tgcacgtggg gtcggaactg cnetctccc ccgggtacca
 420ccgtacncc tccnaccgca aggattacat cccctnaaaa nngaaccctn nctcttgna
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 524

<210> 772<211> 277<212> DNA<213> Homo sapien
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 60ggagggggtc caggacagatg gtctcagggt ggcctcagat gtactcctg aacctgtgca
 120ccagctccag ggaactggcta tcaattctgt cnggtctcaa ggaatagct agtcagctgt
 180tgatgatga cgtccagggg gccctgtccc tgcagtcggc ggctaagggt cagctgca
 240accctacac cctgcttccc gcacagctcg gcgaagg
 277

<210> 773<211> 294<212> DNA<213> Homo sapien
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119

60ttggccctaac accctgtctg actctctccc gctgcagcag ccagtccttc ctgcaactcca
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294

<210> 774<211> 559<212> DNA<213> Homo sapien
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420ccgcantcn agcntgtnc ccaatngcaa aagntatta cmntaaaca aaacccctca
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559

<210> 775<211> 573<212> DNA<213> Homo sapien
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573

<210> 776<211> 592<212> DNA<213> Homo sapien
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592

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120gtgcagatata aaatataaaa aggtttgatt ctgaatagac caactgtcaa ttttctbtaa
180aaaaattttt aatttgggtg agtaaaaacc aaattagttc actgaatctc attttgtagg
240taagagtctt atttgcaata cgaaaaactg agcttatgac tgctttgatt ttctctgtag
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372

<210> 778<211> 381<212> DNA<213> Homo sapien
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120

360gaaggaagc tttttacotg c
381

<210> 779<211> 530<212> DNA<213> Homo sapien
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530

<210> 780<211> 465<212> DNA<213> Homo sapien
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465

<210> 781<211> 378<212> DNA<213> Homo sapien
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120cttgatgcagc gagtggctgc aatgaggtca cncogtgggt ggaatgtaaa gagatcttca
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240gtctccacaa tntggcccca tcaagatgtt ctcatnagt gttaacnagn gcogtctcag
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378

<210> 782<211> 430<212> DNA<213> Homo sapien
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420gcctttttac
430

<210> 783<211> 364<212> DNA<213> Homo sapien
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120agaaggttga gatggggctg cangtgacc cccaaaagga gaagaaatct cttnactcng
180atgagagtga ggaatnaagaa natgacttcc cncnaaagcg ccaagcgctt gaaggyctca
240ttnacatcta gaaccccaac cnggtggmac anacaaacaa aaaggncaac cnaactngatc
300ttngaccggc ccaangagct ttntaggaga gaaccocaaa gagattntta ancncaaacg
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364

<210> 784<211> 442<212> DNA<213> Homo sapien
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120ctctgacaa ctgggacaa ctggggggcc taactcagaa gcaangaa gctctgagc
180ggacagagaa actgctggag accattgacc anctgtacct ggagatncc aagcgcgctg

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 180cctctctctg ggaccactgg gtacaagaga tgggagtctc cgacagcttc tccaatatg
 240aaactaatct taaccctgtg ctgtcagata cctgtttct ggagtcacat cantgaggag
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 180antggatgga cagcttgctc agtaacctgg ggtgccagtc tgctctnatt gtatggncct
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 120agaaatctcg tgctatccgg cnagcccaagg aagaacggac ctccacatc ttctattatc
 180ctctgtctgg ggctggagag caactgaaga cagatctct gttggagccg tacaacaaat
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 300acagacacat ggaggccatg aggattatgg gcataccaga agaggagcaa atgggctctg
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 120aaacacttncg ttacaggaaa tgtatgaocg aaataataa aaattaaaag gtgaaaaata
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 240ctgcagacag cgacaggaaa ncagagcctc ccaactgatt ctgggggacc tggtaataaa
 300aaactcagccca tgatggcgct atggcctctc agacacacca cgctgctaaa acactagag
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 420cgccggcgct cgaaaanggc cgaattncag ccaacttggc ggnccgttac ttntngatn
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 120taactttacag tatcaacatg acgctcttcc tcccttgaca tcaactacat gagaactggg
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 240cagccacacg agtgaaaaga ctgcctctcc tccgncctcg acccagccct tccctgagtt
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 420tgtcacagct acttgaatac accggnctct anatatgcc aagttttaca aatcanacaa
 480aacacaatgg gacacttttt aantgttttc ttatnncca ttatngcctn gnaaggaac
 540ttncocggg ggggncngtt tccaangggg ggaaattcca acacacttgg gggcggttc
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 611

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122

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120gtgaaacaccaa ctatttgtgc tcaacttgcat ctaagtgaag cagccacagc tgtgtgaggt
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300caagtcagtc ttgttcttta tctttatcaa tcaagtctaa aaaaaaaag caaaacccna
360acacacatcc ccaaacata actntcaatc acataggcta attggttcat tattttgmaa
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498

<210> 791<211> 333<212> DNA<213> Homo sapien
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120cactgaggag gatgccasag ctccaaagga ggcctntgaa gacgggaaat tatggaata
180aagtacottg gactgggcca aacctaaagg tgaagggtgc ttcgggggtc ntggtaggag
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333

<210> 792<211> 172<212> DNA<213> Homo sapien
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120tcagtagtca gtcttaccga aacggagccc aactggcatc tatctgagt tt
172

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256

<210> 794<211> 310<212> DNA<213> Homo sapien
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300caattacaa
310

<210> 795<211> 149<212> DNA<213> Homo sapien
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60ctctctgctt gtttggaact tgtaattatt tttttagcag taattaaaga aaaagtcct
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149

<210> 796<211> 579<212> DNA<213> Homo sapien
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120aaagatggga ggaataaana cctgttctgg atccccctc cctccagaa taagagcatg
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579

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120gnttgcctcc ctntgaggn ttgaaccac aaccaaagg tactctntt ttctctggc
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338

<210> 798<211> 140<212> DNA<213> Homo sapien
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120accnagccc caattatgt
140

<210> 799<211> 502<212> DNA<213> Homo sapien
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60ccaaattgag tgcttcatgc ctttagatgt acaggtcgac agagaagatt cccgagagta
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420gcggtcgt tactanttg atcccgatc ggcncnaagg ttngccgtaa tcatggncaa
480taggctggt tnnogtgat cc
502

<210> 800<211> 276<212> DNA<213> Homo sapien
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120gtatctctgc tgcgtgagac caaacagagg tgggggagaa ggggtgcacc cttagcggag
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276

<210> 801<211> 387<212> DNA<213> Homo sapien
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120gtctctctna gattggtttt ttccacnct ctaaaagagt caatcagtt tncacgggggn
180tgcacatagc atttccacac ctctctccaa atcaagtatt tgnngtccct taggancacc
240accgtagata caagtacact tcaagcgaca tgcctctcaa tattcaagca gctacnttgc
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387

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120gtctcgggag acagggttgg ggnctccagg gtctgcacac accccaactc tcttctagc
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360agccctgggt tctgaggcac agtggggcct gggaaacagg caagantctg gtctcaaat
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480ggccggttac ttagtggatc cagcttngga accaanaant gggcantaant cangggtcat
540ta
542

<210> 803<211> 542<212> DNA<213> Homo sapien
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<210> 809<211> 501<212> DNA<213> Homo sapien
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125

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 360attccccaat taacctnnnc aagncaaaag gacctntgg gcngnaaac ccccttaaa
 420gggggaaatt tncacccaca ccttngnggg ccngtntcct tagtgatgat ccacagctgg
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 501

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 120aagaagagcgt tgtagaagtt ttggcaattg ataatacttc agtgcgaaac agcgagattg
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 554

<210> 811<211> 377<212> DNA<213> Homo sapien
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 120cttttcaacg cagcctagt aatgggggag ttggtaatta atgtgtatat tgtaactgaat
 180ttctgtcann taagggggttc actgcttttg tggaaattgg tggaaattgc tagncagtag
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 377

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 511

<210> 813<211> 234<212> DNA<213> Homo sapien
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 234

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 120gctctctgtg acaagaccca agcctggaen gttgncattt aaattggcac canttcttg
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 258

<210> 815<211> 145<212> DNA<213> Homo sapien
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 120ccacggtgc acnaccnntc tctct
 145

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126

<210> 816<211> 231<212> DNA<213> Homo sapien
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 231

<210> 817<211> 238<212> DNA<213> Homo sapien
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 238

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 124

<210> 819<211> 451<212> DNA<213> Homo sapien
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 451

<210> 820<211> 476<212> DNA<213> Homo sapien
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 476

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 466

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127

420anggaagcg gttataatng gntggnggg gngtcnaaaa tggnccttcn ttttttagna
480nacccca
487

<210> 823<211> 525<212> DNA<213> Homo sapien
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525

<210> 824<211> 317<212> DNA<213> Homo sapien
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120aanaangana aatatanaag tgantcattt nccatngaaa aanggcattt ccagcctcaa
180ontaaactca actagttttt attgcattat ttttgaatg ccaagaaact ggccttggac
240ctgcccggcg ggtogctona agggcgcaatt cncncactt ggcggccgtt actnggtgga
300tcnagctcn gnaocta
317

<210> 825<211> 242<212> DNA<213> Homo sapien
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120cttttagtgtt gtgtatggnn atcatttgtt ttgaggttag ttgtattacn cattgttggg
180ngnggattan ccngtttggt catnagatat ttncangngg ggatcaatc agggggaaat
240ac
242

<210> 826<211> 348<212> DNA<213> Homo sapien
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60tcaccgcgag cactttaatc cccacaacag cctgtgagg taggaatcac cacactccc
120ccnactcat attacagatg ggggaaacn agacacacat ttncaaaagc gcgcacaca
180tggggagtto anggttagcc tgnagtacga acacacttct tocccatggn ccagtcagca
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348

<210> 827<211> 349<212> DNA<213> Homo sapien
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60tgaaaaatgc ttttccctc octcacagca ccgttttata tatagcagag aataatgaag
120agattgtctag tctagatggg gcantcttca aattcacoca agacgcacag tgnntatttt
180accctccctc octcataaga ctttaaaaaa aagaaaaaa caccctnca aaaaaantca
240aanaatttga ggaaccocct ccaaacagtn cacagttatt agttcangt ggtcaataat
300tcacatcttg cancaaaagng tatggacatg atttcttttt caaaacttt
349

<210> 828<211> 191<212> DNA<213> Homo sapien
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120tataaatcta ggtcttctgg gtcattaaan gtattaaagt tcaagtgnctt ttttttttt
180tnngccctaa a
191

<210> 829<211> 447<212> DNA<213> Homo sapien
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60tcaactggag tccgtggcta tctctccga gctgtttatg atcagacaga ctggggagac
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180aaactggatct ggcgcttcta ctttgagggc ttctttgacc toattgtcgtt ggtggccggc
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300ggaaagaagc tcaagttggc cngcataagt gccaaanaac atacaccaga tctgtctctc
360agggtgtcgc gacagaattc ttaccacago aaaaacataa gatgttgat acngaaaatc
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447

<210> 830<211> 548<212> DNA<213> Homo sapien
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60aaacttaactt taataaaggat ggctgcaaag atggnaaagtc ttaactgggtt ttaactgttaa
120ctctatttctt ggacataact atgaattttt tatacaatgc acttcatgaa aagtgtgggc
180tccccccagat tgcccacaaag ttgtgatcttg aagtcctaaa catttgtcca tghtaagcttc
240aaaaaacaggt taactgagtt attcaagtag caagtactta aagatacaat tcttgaaagca
300gttttcaatgg tttctgatcc aaataaatcag tttctgacat taactactca cataatagaa
360gcatctctca gtttcttctt cactttctct tttctctttt ggttnccctt ttgtggncctg
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480ataanggttg gggattgacn tgcctacctt ggcenncnct taactcttaa aaatcaaatg
540anaaaggg
548

<210> 831<211> 183<212> DNA<213> Homo sapien
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60aagatagaa tggtggttgg ggttctgggc agccactgct tcagccctgc caagctgagt
120gtaccgagca tgagactgtg aggtacgggc cccataccat ggtgtcaaca taactctgga
180agg
183

<210> 832<211> 169<212> DNA<213> Homo sapien
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169

<210> 833<211> 351<212> DNA<213> Homo sapien
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120catagtctgt gctgactgtc tctgggttat agttcaacct gatgtgtcta tatccactc
180ttcggagctg ctggatgca octacanac accagtcaaa tcaacogta gagccataac
240ggtagagccc agagccaag actaggacat gaggtgttgc aaagtgagg tcatgggttg
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351

<210> 834<211> 478<212> DNA<213> Homo sapien
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120cagttaccaa agcctagata cgcgttagat ggcctcttcc cggctgtgac gttgtctgt
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300gttttgccaa atccactctt ggaacccggt ccaacctatt tgcaagtagg ttgtgacctt
360gatgaaactg catctctact gcaatgag gcttttatt tgtanggaca agaanganga
420gtttccgntt atttttgtaa cntgntttac attggttcgc atntaagtna aattcggg
478

<210> 835<211> 421<212> DNA<213> Homo sapien
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60cctggctcgg gaagggaaga gaaaaaagac gcaggccacc tgggggttct cgagtctttg
120gtcagtcag cttctctatc tagctgncct ttgcttcgcc agtgtaaaac ttgcctgcgc
180ggagtgagga ggcacagctg gaactccnag ggcctagag aggcagacag catcttggn
240tcaagcttgc cttctccttg agtccctctc tccctcnag tctagcaga ggtgtagct
300gcagatctat gaaganaaga actgggggag gaggtagag gaactcggc ggcacaccc
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420t

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421

<210> 836<211> 515<212> DNA<213> Homo sapien
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 120aaggtccctga aactctctg tgcagctctt ggatccacct tcaatagtta tgcctatgac
 180tgggtccgco aggtctccag caagggctga ggtgggtgg caattatagt atttcttggg
 240aataataaat actatgcaga ctccgtgaag ggcgattca caatctccag agacaatna
 300aanaacacgc ttgtctctgc aatgaacag cctganaacg tgaagacaac ggtgtgttat
 360tactttntgco ganacacnng ggggtcnatt gctttntga atggtntctg gggggcccaa
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 515

<210> 837<211> 416<212> DNA<213> Homo sapien
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 120ctggtgaacg anaagtctcc ggacatgac catttcttgt ggtgggcagt gaccaccatt
 180accaggtcaa cggtaanaag gattctttg aggaagacca aagggggggn accatccaan
 240ttgaaataca cncnccactg tcgagtttgc ctacctggcg gacctntca tnanagcng
 300acatncmaaa cattcaaan agcatacna ctantcttc acttttaang cctactcnn
 360ttgnaaacn ccttaactat gggcntngtn ttcttttga cctngtnc tttatcc
 416

<210> 838<211> 58<212> DNA<213> Homo sapien
 ctgantcaag ccaaaaaaaa aaaaacccaa ancaannaaa aaanccantt aagncctt 58

<210> 839<211> 193<212> DNA<213> Homo sapien
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 60ggtttgtgcg aaaaattatg tgaacccgog gagacccttc gagaatctc gtctcgacca
 120aaaantgaa ncttgatcgg tgagtatggg ctccggaaca aacgtgaggt ctggagggtc
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 193

<210> 840<211> 468<212> DNA<213> Homo sapien
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 60caattccatg agggaaaaaa actacgggag gaaattctac aacaccattg ctgccaccac
 120ctgcgaaggnn cagctttctc actagtagtg aaagaagcg tttctgagga caaattcaac
 180ttatgtacaaa aaattgatac agccatttcc aaagagacga gtaattgaca caatggcagt
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 300ngcgtcgaga gccanaaagg gggcaacgog aagacnagtt tntagcgacc cttyggaaaa
 360gctacagcta cacatttccg agggagattaa aacattccat atgccattta actttaacct
 420aaaaaaaata tagtgggagg gaaccttgn ntcngagaa attcaag
 468

<210> 841<211> 449<212> DNA<213> Homo sapien
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 120ctggggcctt gtggnacnac acagccagta ggggttaggg ctgaagacca gggcgttgat
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 300canttgaggg acttggagac agctctctat gatttttgt gaaggacct ggagcgagcc
 360accttaagg cgaattccan nacactcgcn gcggttacta attggtatcc aantcgggtc
 420caacctttgc gnaatcatng gtcantct
 449

<210> 842<211> 177<212> DNA<213> Homo sapien
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 60gctcctatgt cttyggccca gctcgcaccc caaagcctgg ggaagagac ttttccatg
 120ngcttgaaaa aaaaatgcgc ttggagatct ccaagcctc ctggttactg tggctgg
 177

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<210> 843<211> 123<212> DNA<213> Homo sapien
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 60gcaggagacan ggagctgggt ggggaggacc anaaatcagg ttatcaatac ttttggntga
 120cca
 123

<210> 844<211> 507<212> DNA<213> Homo sapien
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 60ccttcagggg tacacgcttc cccatatgac ctgtgaagac ctcaagcaac tggaaatgggt
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 300gagagggtgga gctcagagga tccacagntg gatagatgcc cagctcagca atggcacgag
 360acntacatt ggtttgcattc caaatgggca aacctntnta ncanggggca ggggtcangt
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 507

<210> 845<211> 434<212> DNA<213> Homo sapien
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 120atagcggcgt caccatcggg atgtctctgat ccaacatcna ggtcgtaaac cctattgttg
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 240gttattggat caattgagta tagtagtncg ctttgactgg tgaatttata gcatgtgaat
 300gctcggagagn tgggttctgc tccgaggtgc cccaccgaa atttttaatt gnaggttttg
 360annttttagg acctgtgggg tttngtaggn acntggctcg catnttatac gattaananc
 420tccantnggg gctt
 434

<210> 846<211> 317<212> DNA<213> Homo sapien
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 60tgtataagtt tctgtgtgct gatcaaaaga agaaggcata cncagatttc tacagaaact
 120acnattgctm tgaaaagattt tgaggagatg aagggaagct ggtatcttcc anagtgtaaa
 180gtaattcttg aatataana atttcttcag gntgaattac ctanaagttt tgtcactgac
 240cgtgttctcc gaactatgac acatgaatat gtgggctaag aaatantcct tcttgataaa
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 317

<210> 847<211> 464<212> DNA<213> Homo sapien
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 60ctctangaga gcaaacacat gttctacttc tgtatgtccc tccctcattt caaatgagaa
 120gtacccaatt tngtataaaa taaccaaata accattgccc caccatgaac atggggcttg
 180ggaagacagt cctacaattct tcatcatata tttaggtttt taggccancc agctcttttt
 240tctccaaagct ttcttttgaa tgttccagat ctattttaat cctaactata gactactgty
 300ntttgtgagg tgtctgantg tctatgtgag ggcaaggaca acantgcagt ccaataaaca
 360cnaaaaaaat tgcctttttt gcanctgaag ctctgntctg ganatttcat tttgttact
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 464

<210> 848<211> 561<212> DNA<213> Homo sapien
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 60attttattgc gaaggagctc gtgaactgta gttcttattc ttgggtccaa aatcagaggc
 120tgtattacaa aggaagaaaa gagtgtattc aggggagcaa ctagcttctc gtcacagaaa
 180tccaagggca gagggtgttc tgaaaggggg gatgtgtcag tatcacaana acaatcaaga
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 420tgagggtggt tggaccocag agtgcaaatc caggccccag tgacagtaca ccgncnggaa
 480tggaatgag ggattaaagg gctntanaaa accaaaagga ctgctgttaa aggcagagat
 540tgannangaa ggagaccccc t
 561

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<210> 849<211> 428<212> DNA<213> Homo sapien
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 60gcaagtgactt ggaagtgctt atctcaatgg ccccgccacc agccaccact gaatcattct
 120tgatggncoc cctcgacnat catgatggca tcatgcaggg accgctctgt cctctccata
 180aactgtctcgg cgcgcgcaac gagaatgaag gtccatgtct tggccttggg gcagccagta
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 300acatctgctg acagagcatt cacactggtc tggattgagc ctccacaggc catcattgtc
 360ctcttcagat cctctcagg tactcggcca gcacagaaca tgttctctgc agcaaatgac
 420tgggtggg
 428

<210> 850<211> 391<212> DNA<213> Homo sapien
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 60ctctattttt ctcagatatt gtaagcatto tgtttttcaa tattgtagtt aattttttgg
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 360ccctctctct tggacctcgg cccgcgacca c
 391

<210> 851<211> 329<212> DNA<213> Homo sapien
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 60tncaaagcct gaacgacgc ctggcctctt acctggacag agtgangago ctggagacot
 120agaacccagay gcttggaaana gcaaaaatccg ggagcactg nanaagaagg gaccocaggct
 180oaganactgg agccattact tnaagatcat cgangacctg agggctcaga tottngcaaa
 240tactgtggac aatgcocgna tngttctgca gattgacaat gcccgntttg ctncotgatga
 300ctttanagtn aagtntgaga cagagctgg
 329

<210> 852<211> 279<212> DNA<213> Homo sapien
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 279

<210> 853<211> 267<212> DNA<213> Homo sapien
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 60gaggttagtt gtggcaataa aatgatttaa gatactagt ataagagato aggttcgtcc
 120tttagtgtttt ggggatttgn aattatttgt tttagagtta gtttgattag tcaattgttg
 180gtgggtgatg gtccgttgtt gatgatgat ttggagggtg ggaatcaatg agggggaaat
 240ngaagatgat ctactgcngc gggatg
 267

<210> 854<211> 335<212> DNA<213> Homo sapien
 aaaagctctt ggtaggatta gttggttcta aggtacocct tagggacotc attatttcaa
 60gaggaaacca aagtccagcc tctacatag atgctgcoco acgaaggacc caccaaaacta
 120acctatttt aggtttctca ggnangcagt tctgtctcag cttagagcag aaccataaaa
 180atactcaagt actgggatag gcaaaagcat tgtgttact gtggatttgt cctggaagc
 240ctcctttgggt gagaacatgt gaaccaggca cctgtgtttg tttagagcat tgcgtccacc
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 335

<210> 855<211> 348<212> DNA<213> Homo sapien
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 60acaggcaact gccaggttgg ctcaacatca cgcacatcg tactgagact caaggccgtc
 120tccacaactt ccaaccagtg caatgaact agtgcaaaa aattcagaa ggaagcgggg
 180aaacagagtc gtggagctct tgaatctctc agaaaaagg aaagcacaga aagctcagaa
 240acaagaagac agaaggatga aaaaagaana gaggagagtg gtggggagcg cgtatcccg
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348

<210> 856<211> 371<212> DNA<213> Homo sapien
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60tggcggagtt cctccacggc taaggcagct gtgacgagat cccagggtg gacttcggg
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240aaacaagacc actgaanta cccattntn gggaganagg ggaacaatn ttoaccnaca
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371

<210> 857<211> 358<212> DNA<213> Homo sapien
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240actggggnac gtacatggcc ttgtagcacg aaggnocact ccaagggtct gganctgccc
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358

<210> 858<211> 346<212> DNA<213> Homo sapien
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60ggaagtccaa ctacttccct aagatcatcc aactattgga tgattatccg aaatgttca
120ttgtgggagc agaaacaatgt gggctccaag cagatgcanc agatccgat gtcccttnc
180gggaagctgt tgggtctgat gggcaagaac accatgatgc gcaaggccat ncgagggcac
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346

<210> 859<211> 380<212> DNA<213> Homo sapien
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60ttcacatctg tggcacagag ggcaacgaag gtctgcgctg gtttcggat gatctgttg
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380

<210> 860<211> 328<212> DNA<213> Homo sapien
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180agagacttga gccattactt caagatcagc gagacactga ggcctcagat ctgcgcaat
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328

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180gggaagctgt tgggtctgat gggcaanaac accatgatgc gcaaggccat ccgagggcac
240ctggaaaaaa acccagctct ggagaaactg ctgcctcata tccgggggaa ttgtggcttt
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346

<210> 862<211> 209<212> DNA<213> Homo sapien
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120tgnaggctgc nattgagcaa gatccnccac ntgcactcca gctcgggcaa caagagcgaa

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133

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209

<210> 863<211> 328<212> DNA<213> Homo sapien
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328

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563

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120tttggtgtgaa atgtggagg ttatggcaaa tacagggtag gctgaatggt tctaggccag
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240gagtaacccg ctgactaatg atccacatg agagactcaa atctaacact tggcacaatt
300acaactctggg atactgaggt cccactcttt agtgaagcc aaaggcccta nggtcaaaat
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420anaaaacact atcacccctc ccttcaagtt ttaatgaca acnaggaaac tggggctctc
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538

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120gatgtgaaat ggcagtcact taaagacctg gntaaagaaa aagctgggtga ggtacacac
180gtggagctct taatggagcg tgaaggaaag tcaagggtat gtgctgtgtg tgaattcag
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300ccacttgaaa gtcaagaaga atcctgatgg tgaacattgc cngngagagc aatgccaaaa
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420gaaacngaa ggaagnnttt nctctggct gggggngng gctccanaca gaentttttg
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534

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120gtaaactctt cttctctctc cagctggcgg cccacgctca actgatagtt acttgatcca
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295

<210> 868<211> 461<212> DNA<213> Homo sapien
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120aaagccatgg ctctgaaagg nggcaggcca gaaggaaacc tccgttcanc taaaagtga
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134

240accagatct tccagtgatg cactgtctgc ctcttttaat ggagaaatgc tgggggaaccg
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 360tcataatnccc ancccttnat ntctgcctat tccagcttna nggatgttna ctngagccccc
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 461

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 180tcagtctgtg cctttttaag agaacgcat gtctctact ctgattctat caaaatgtgt
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 519

<210> 870<211> 161<212> DNA<213> Homo sapien
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 120cagctccat cccgagggg tatcaccgcc gatcacatag g
 161

<210> 871<211> 536<212> DNA<213> Homo sapien
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 536

<210> 872<211> 327<212> DNA<213> Homo sapien
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 120cctgggtggc aagacagga cctctccca gntctccata cctaagggg tcatgacag
 180caagggtagg accmngccc ccagctcag tggtaagg gggncctcac ggantcccc
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 327

<210> 873<211> 446<212> DNA<213> Homo sapien
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 180aaaaaaccc aaaaagtaaa aacaaaaaaa acaaaaaacc caagaaaaaa aaanagtca
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 300naggagacc tgnrtgntna cctggggcca caaanccctg ntttgtccc accattgcag
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 446

<210> 874<211> 302<212> DNA<213> Homo sapien
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 120cctgtctctgt ccagcttcag tgacacaagc tgccttagct aaagtcocgc gggctcoggc
 180atggctagc tgagagcagg gatctaacgt gcttctcagt tctttgtgtg gaaggagac
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135

300gg
302

<210> 875<211> 374<212> DNA<213> Homo sapien
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240ttggaagat gtttactct cgggaatctt ctgttcaag ctgtacatct aaagcatga
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<210> 877<211> 538<212> DNA<213> Homo sapien
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<210> 879<211> 231<212> DNA<213> Homo sapien
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60aaagaataa gaaggtgtaa aaaaaaattt ttcaaacccc aaataatgat aaaaatgat
120gtatcctctg taataaatct ggaactaaact atcagctcat tcatggttat tcatgttaca
180gagcatgaag tgaacaacca aagtataaaa aattaaaaaa aaaaaaang a

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60caaatatttt ttgcatttat gaacggctgt ttttctaatt tgaattgtg agacatttc
120ttggggaggg aaatttgaaa ntgggttccc ttttttagaa attgaagtgg tottcattg
180tcaactacag aaaaggaaaa aaatagaatt tgaaggattt ttatgaattt atattgcatt
240actattttga gtcaaaactt gatccttgtt ttggaatcca ttgtcaatt cggaatgaaa
300aattataatg taattttaca ttacataagt cccattttaca attaaaaaat agcaactctt
360catcttatgc ctgttgagaa gatattaaat tttcacattg nngacagtga aatgctatgt
420ttgggtttata aagantacmg accat

<210> 881<211> 414<212> DNA<213> Homo sapien
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136

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 414

<210> 882<211> 554<212> DNA<213> Homo sapien
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 554

<210> 883<211> 108<212> DNA<213> Homo sapien
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108

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 300t
 301

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 136

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 399

<210> 887<211> 326<212> DNA<213> Homo sapien
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 326

<210> 888<211> 531<212> DNA<213> Homo sapien
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137

120tgccattaag ggtgtgggccc cgaagatatg ctcattgttg tgttgaggaa agcagacatt
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 420ntgncocgagg gccanchnaca nccaaaaancc cttgngccgc ccgnggcggg acccggtggg
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 531

<210> 889<212> 581<212> DNA<213> Homo sapien
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 180

<210> 891<211> 124<212> DNA<213> Homo sapien
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 120ngaa
 124

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87

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 420

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 60gttggaacag atctactgtg taaagcaaaa tgaaccaatt cttgaaaagt gttccagggc
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 180aaagttaatgg aattaaaata atgataactg tagacctctc tccctaanga tgggtgctg
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 314

<210> 895<211> 353<212> DNA<213> Homo sapien
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240ctctgagagct ccaagggagt ggcccagccc ccattctctt gactttagcc ttctgaaaa
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353

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120agaagaccca ttaagaggtc ttctgggagc cttaacannn ccccatattn cccocnccag
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240ggtttgatga actcagaggt acaatccctg catacagaat gatttnggct nttttmaag
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420tgatggnggc cccac
435

<210> 897<211> 331<212> DNA<213> Homo sapien
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120ctctgcccac tgcgtgtgtg aaggacctca gtccaagaag cgttcagact ttccagaagg
180aacgagccat ctctcgtggt gctcagaagg aggcagattt ggctgcccaa gaagaagctg
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300gtgcccnang ggangaagggn gggcncocca a
331

<210> 898<211> 690<212> DNA<213> Homo sapien
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120cagggaggaa ttgaaagtag atagaaacgg aactggatta ctccggtctg aactcagatc
180acgttaggact ttaactgttg aacaaacgaa cctttaatag cccctccccc tccggatgct
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360gttcgctttg actggtgaag tcttagcatg tactgctcgg aaggtggggg ctggtccnaa
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480actgtttgnt aataaattaa actccatagg tntctgctgn tnggtatgcc cnettttcgg
540gcaggcaatt tactgttaaa gtanaaanag tgacctctga acctatcag ccnattagg
600acanggatat gtctctcccc cgcncgcocn cgcngttnact ttatnngng ggnntttac
660ngnagtaga gggagnttgg taannggggt
690

<210> 899<211> 432<212> DNA<213> Homo sapien
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120ttgactttat ggagaattt tcaactggaag gaaagactaa cttcttttg agagagatg
180cgtagatata gaggatggga gtaggtgcaa gtccaaacaga gaattctttt accttgatg
240ctgacttcta aatgaactga agatgtgcgc ttacttggtc gatttttttt ttccatctca
300taanaaaaa caagtgaag ggtcccaact ntcccccccc ngaaatggcc cgnagtttt
360taacaaantt ttntcttcgg gggggccctc aaangngaa ttccncccn gggggcngtc
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432

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120tcccaactc caaccagtgc aaatgactta gtgcaaat taattcagaag ggaagcggga
180aacagagctg tggaggcttt gaactctca gaaaaagga aagacagaa agctcagaaa
240caagagaca gaaggatgaa aaagaagaag aaggaggttg tggggaccgc gttattcoct
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378

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438

<210> 902<211> 327<212> DNA<213> Homo sapien
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120tgagcgggtt acttatittg acagatatca ctttgggtct otttaacatta aatttcttt
180ctctaagtaa tataagacat acccatagc tctgtgtgag ccagcaaatc cgtgcocccc
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327

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120aacccagcaac cgacctagag aatcacggaa gtctcttatc aaagcagata tgtgtcaatg
180ctgtctaaac acacacattc ttttctgggt gccccnattg taattattt nangagctgg
240gggacnnga cacgtttatc ta
262

<210> 904<211> 482<212> DNA<213> Homo sapien
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120tcacacaactc caaccagtgc aaatgactta gtgcataata aatcagaag ggacggggga
180aacaagaatg tggaggcttt gaattctctc gaaaaaagg aaagcaggaa agctcagaaa
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300gaaganccta antnttggat taattgttgg gnttgggggn naaccctccc cccaggggng
360gactgcccgc cggccttnaa ggggaattca nnncttggc gogttattnn ggatccaact
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480cc
482

<210> 905<211> 224<212> DNA<213> Homo sapien
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60ggtattgggg ataataattca tttagccttc tgagctttct gggcagactt ggtgacctg
120ccagctccag cagcctctct gtccactgct ttgatgacac ccacgcgaac tgtctgtct
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224

<210> 906<211> 326<212> DNA<213> Homo sapien
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326

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369

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211

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331

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240agactttgaa gaaccttttg gatgtggggc atcatccga tctttctctn tctccaaaat
300gacaaangtt ggggaatttt ttaat
325

<210> 911<211> 313<212> DNA<213> Homo sapien
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313

<210> 912<211> 360<212> DNA<213> Homo sapien
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360

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300gacttttaact ccggnognga ccaccttaag gnaattttc ncaactnngg ggcgttntta
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415

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314

<210> 915<211> 403<212> DNA<213> Homo sapien
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 360tggggggngg tttagggganc cancttngnc caaattggg aaa
 403

<210> 916<211> 83<212> DNA<213> Homo sapien
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83

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 347

<210> 918<211> 339<212> DNA<213> Homo sapien
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 60cttcaggcac ctgctgtgccc tctctctcog cagatgcctc ggttggaagc ctccgtcact
 120gcctctctgta acagcaccag ctggacgttg tcatgaaatg tcaacagttc tgggtgtttc
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 339

<210> 919<211> 102<212> DNA<213> Homo sapien
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102

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 120ccagagatcc aggaacggtg tcatggagcc cacttctcga cagactgat gatgtggagg
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 504

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 447

<210> 922<211> 375<212> DNA<213> Homo sapien

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 375

<210> 923<211> 479<212> DNA<213> Homo sapien
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 479

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 576

<210> 925<211> 321<212> DNA<213> Homo sapien
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 321

<210> 926<211> 348<212> DNA<213> Homo sapien
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 348

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 180caactcgtgaag atcctaaacc ttcttggaaa tgaattgagt cttgagcggg aattggacaa
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 319

<210> 928<211> 335<212> DNA<213> Homo sapien
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60ttgggggaaat catcatctgt ttctaaacga aagctgcagc ggaatgagag tgaaccttca
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 180agtfgaggagt ctcttacatc tctccatgca gtgtatggtg attctaagct ctctgaccca
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 335

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301

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314

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295

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 612

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462

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396

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480gttacctgcc cnggcggccc tcgaaagggc gaattccaca cacttggggc cgtntcaang
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582

<210> 962<211> 114<212> DNA<213> Homo sapien
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114

<210> 963<211> 601<212> DNA<213> Homo sapien
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480gaagaagcnc atgttaaatg atagctctta atgaggaat atggcgtggt actattctct
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600a
601

<210> 964<211> 560<212> DNA<213> Homo sapien
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560

<210> 965<211> 223<212> DNA<213> Homo sapien
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180caggaggggcc gggcttttcc tctccctggt cactgggagc tgg
223

<210> 966<211> 425<212> DNA<213> Homo sapien
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120ggggaaaatg ctattctgtg ttttggaaa gaagaaatag tgcctgccta ttatttcta
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300caagagctta caaaaaactaa gtggcattgt attttataa cccattgag aagactaagt
360aagaatgaa atgtcctatc aattttattt tgtcatgctt caaacaataa agacatttct
420gcttt
425

<210> 967<211> 339<212> DNA<213> Homo sapien
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60tcacgtgtgc cctgcgatgg gtcccgtgtc actgtttaca tgacctattt gtgtggttat
120atagcccttt atttaaaaga gagaagtctc ttttacaagg ttattaaatt aattatagt
180ttaaaagtta aagaaaaaag agctgcagag tattttataa actgtctttt agaaaaaac
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339

<210> 968<211> 291<212> DNA<213> Homo sapien
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120agtggagaagt caccgcacca ccacgtccct ccgttctctg ttggaccccc ccatcctacc
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291

<210> 969<211> 130<212> DNA<213> Homo sapien
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120tgccccaga
130

<210> 970<211> 210<212> DNA<213> Homo sapien
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210

<210> 971<211> 122<212> DNA<213> Homo sapien
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60aacctttgccc cttcacaata tagaaaatat tggttttgccc attacatttt aatgcacagt
120tt
122

<210> 972<211> 108<212> DNA<213> Homo sapien
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108

<210> 973<211> 313<212> DNA<213> Homo sapien
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120gtgtgcagga ctgctgtgaa nactgtatgg acnacgggn agttcaantc ctttttctat
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240atacaataac cactgaaaat acccantgtt nggtagacga ggggaacatc tcancacat
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313

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<210> 974<211> 272<212> DNA<213> Homo sapien
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 120gttntaagt ccacacatga acaaatana accttaataa aaggtcagtg ttaatgcaa
 180tactagcata ggttcagcac caagcncaat gttattttac tggttngcot ttttcattct
 240gtttttttt ttttgttttg ttttgttttt tt
 272

<210> 975<211> 375<212> DNA<213> Homo sapien
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 120tttggagg gagccatcaa aactcgatca ggaatgtttg gcgcgtcttg aatgtcaga
 180gtcgaccc cactcagttcc cgccgtgca cagatggaa agtgcgtctc tgtgtactc
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 375

<210> 976<211> 340<212> DNA<213> Homo sapien
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 60cctggtcttg gaagggaaga gaaaaaagc gcaggccacc tgggggttct gcagttcttg
 120tgaagtcagc ctttttatct tagctgcctt tggcttcgag agtgaacc ttgcgtgcc
 180ggaggcagga ggcccagctg gacctccagc ggccatgagc aggcagcagc catcttgccc
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 340

<210> 977<211> 429<212> DNA<213> Homo sapien
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 429

<210> 978<211> 390<212> DNA<213> Homo sapien
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 60gtatgcaggt tccaactctg tgcagagctg catcttctatt tacaagtctc tggtagcact
 120ttanaagtga agcttggctt caaaatcaca acactggggg ctttggtctca accctttaa
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 240tctgactgaa agtccctctg agtgcactct tnggtgcac atggccgccg cacacacaa
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 390

<210> 979<211> 372<212> DNA<213> Homo sapien
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 180aagccacact agcccaagtt gtacgtgaat gtttaatgt cttcacaac atggaaaata
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120tcaaatgttt tgaatttttg aacatttctt gtttctgact tttagattag ggatactcac
180cccgtaacca tatttggagt ccttagacat tagattatat gaaaatgatt gattgatagg
240taagaaagtt tacctggccc g
261

<210> 981<211> 266<212> DNA<213> Homo sapien
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266

<210> 982<211> 199<212> DNA<213> Homo sapien
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120gaataggtgt gtgtagcgac actagtgaag cgagtctgc tgaattngat gataggcngg
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199

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344

<210> 984<211> 400<212> DNA<213> Homo sapien
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400

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232

<210> 986<211> 347<212> DNA<213> Homo sapien
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347

<210> 987<211> 439<212> DNA<213> Homo sapien
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360ttctctctggn nnaccactgg cntctntagat ntnnagcntg gtggccancic cgnacccttn
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439

<210> 988<211> 256<212> DNA<213> Homo sapien
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256

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380

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366

<210> 991<211> 302<212> DNA<213> Homo sapien
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300cc
302

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569

<210> 993<211> 362<212> DNA<213> Homo sapien
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360ga

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362

<210> 994<211> 501<212> DNA<213> Homo sapien
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501

<210> 995<211> 374<212> DNA<213> Homo sapien
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374

<210> 996<211> 304<212> DNA<213> Homo sapien
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300ggtt
304

<210> 997<211> 344<212> DNA<213> Homo sapien
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344

<210> 998<211> 542<212> DNA<213> Homo sapien
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180ctttagggtt ggcctcgagg gtnggtggga ttggagggg aaggggggnc ttggnatgan
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285

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<210> 1002<211> 273<212> DNA<213> Homo sapien
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 436

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 180atgacaaagg gcatcccttc caagtgacc accagttcca ggggactatg cccagtagct
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300ctgccaacaa aggcattggtg ctttggagcc cagtcttccc ttggagtctg taccccacca
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<210> 1007<211> 116<212> DNA<213> Homo sapien
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<210> 1008<211> 220<212> DNA<213> Homo sapien
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 220

<210> 1009<211> 96<212> DNA<213> Homo sapien
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 60ataccaagg caccacacac caccgttcca aaaagg 96

<210> 1010<211> 550<212> DNA<213> Homo sapien
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 120actgtggagaa tattgtaggg gataagtctc tgagaacaaa catcagactg aaggcaatct
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 420gcattttgcac tgtngtgcct tnaangaaag gaaaaaaa gtnccactnt tccagannat
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<210> 1011<211> 334<212> DNA<213> Homo sapien
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 334

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<210> 1013<211> 434<212> DNA<213> Homo sapien
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 120aatcatcttt ccaatccaga ggaacaacga tgtctctctg ccaagatcca totaaactgg
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 434

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 120taccttcacg tatcaacatg aacgtcttcc tccctgacat cactcacctg agaactggcc
 180ttcaccaatc ccagagaccg tgcgtaaac acatcaagac agaactgttt gccatttcca
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300ccagttatatt cagctcacac cagacgcgag ctccagaggt gaacaatat ttcataaac
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 420agctactgaa tacacccggat ctanactgac ccantttctca aatcanacag nacgaatgga
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<210> 1015<211> 344<212> DNA<213> Homo sapien
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 120ctcaacgaag gcaaacacct ttacacgcta gatgggtggg acatcatcaa ggcctgtgac
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 344

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 120acttttgccc tgcaccacatt gcaggtgttt tgtatatata caatggataa attttaagtg
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<210> 1017<211> 250<212> DNA<213> Homo sapien
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 120tganattgtt tgggntactg ctacacagtc gcccgatcac ggcngntnnt tgagtttgat
 180gctcacccctg atcagaggat ngagttaaag gntngcctat angcngntat aataaatatg
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 280

<210> 1020<211> 365<212> DNA<213> Homo sapien
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 365

<210> 1021<211> 425<212> DNA<213> Homo sapien

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425

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131

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213

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303

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490

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356

<210> 1027<211> 425<212> DNA<213> Homo sapien
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425

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 577

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 331

<210> 1030<211> 201<212> DNA<213> Homo sapien
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<210> 1031<211> 192<212> DNA<213> Homo sapien
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 427

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 193

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527

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240ctgacttctca aatgaactga agatgtgccc ttacttggct gatttttttt tcaatctcat
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428

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179

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148

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275

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338

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220

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205

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243

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156

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353

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512

<210> 1070<211> 108<212> DNA<213> Homo sapien
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108

<210> 1071<211> 507<212> DNA<213> Homo sapien
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180ggttaaaattt tgcattgtag atgatgaga attaagagag tactgtactc aagccttga
240atcatttgta gaaagatgat ctaaggaaat atactccatg ttcttcaaca ttataaatat
300ttgtctttaa tatcttaact atgatccaaa ttataattac gatgatgaag atgaagaatc
360aaatgcaatg gatgcttgat ggtgggtgat atgatgatga agggagtgat gatgaatac
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507

<210> 1072<211> 377<212> DNA<213> Homo sapien
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120gtccaccacca agctcagagc ctggaagccc agatagacaa gcagagatgc tccagaattt
180aacccccact ccatctatg ctgtacaaat tgcttctaaa ctggcaatct acaatccaaa
240ttttaccacc accctgccag ttaactcaaa aaacatccaa cctgtcagat acaatagaag
300gagtaacccc gatttggaga aacgacgat ncactactgc gattaccctg gntgcacaaa
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377

<210> 1073<211> 359<212> DNA<213> Homo sapien
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180agttcatgtg gttaaaggtc agaacgtgg ccccttacag agctgaagtg ctcccacact
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 359

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 240cgtgaaaaaa gcaagaaaaa gaaggaaan gaggaagat aggaagatga agaggtgag
 300gaggagaggg aagatgaaga anatgaagat gaanaaagag atgatgatga tgaataacct
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 427

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 120cagatgtgct ggttcgatgc caggtgtttg aagagaccga gattggaggc gagaggtaca
 180atttttttac tggctgcgcc aaggccaaaga catgcacact catttccgtt ggcggcgccg
 240agcagtttat ggaggagaca gagcgggtccc tgcgatgacg catcatgac tgcaggaggg
 300ccatcaagaa tgattoantg gtggntgggt gcggggccat tgagatggaa ctctccaagt
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 433

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 240ctcttgagtg agtcttctgt aaacagtcga ccaatgctct atgttatgga aaccagcta
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 420aatcaaatgt ttacttttct tgaacctata ggaacctcgn cgggaccacg ctaaggcgga
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 534

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 420gcaccacaaga ttcagcgtct tgttactcca cgtgtctcgc aacacaaacg gngngcgtat
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 537

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120gtactgtgac agogaaacatc tgalatatg aaaaactgcat catcaattca acgttttggg
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246

<210> 1080<211> 220<212> DNA<213> Homo sapien
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220

<210> 1081<211> 253<212> DNA<213> Homo sapien
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253

<210> 1082<211> 223<212> DNA<213> Homo sapien
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120tgtgatttta tgatacgtat acattgggct ctgtccaagg ctccgtgcctc atgactccca
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223

<210> 1083<211> 534<212> DNA<213> Homo sapien
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120aacctggcac tcaagcactt tgcaacgatgt ctcaaccaac atctgacatc tttccctggg
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534

<210> 1084<211> 199<212> DNA<213> Homo sapien
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199

<210> 1085<211> 469<212> DNA<213> Homo sapien
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469

<210> 1086<211> 199<212> DNA<213> Homo sapien
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 323

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 378

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 426
 426

<210> 1091<211> 320<212> DNA<213> Homo sapien
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 240ctccccctgc accgtcaaca cneccacctc gtccgntgta ctccnaccca tgtttccagg
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 320

<210> 1092<211> 522<212> DNA<213> Homo sapien
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 120cggtgagcct ggcaaggcat tctcatcaac actcgtgttt gcaaaaggtt aaacaaaaac
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420ccoccttaag ggcgaatttc cancacactt nggnggccng tttcttaagg ggcacccnac
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522

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453

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414

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546

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543

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 470

<210> 1099<211> 409<212> DNA<213> Homo sapien
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 313

<210> 1101<211> 306<212> DNA<213> Homo sapien
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 306

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 120ctaagaagt ggcctggaga tgttagaag gttaaaaoca accagaaga aatctaagtga
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 300cttttagttat attcaactgt ttacacaga gaaatacaaa ataaagatca cacatcaaga
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 120ctcaggggaa gagtgctgt tttaaaacgg catcctact acacatatat cccctctccc

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170

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 420ggagcagacag aaggaggcac agaatgaggc acatcctnfc ggcacacctg gnnccnigtct
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 120gtgggaagaa gttgcacatg gagccagagg agtttgactc tgacacccctc ccgcagtttg
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<210> 1106<211> 538<212> DNA<213> Homo sapien
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 180gttttgcaat gaacagagat tggaaaggag taaggactct ctccctctct attcaacatc
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 120agotgatatt tcaagtcccc cagttctgtc cagacaatct gtcttaacota tagcaattaa
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 420ctcccatgg ctaggggaaa atggcaacant tgggtcctat ttaaatgctt aaagaaaacc
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 360aattttccagc tgaataaaaa aaaaatttta agtgttctcg ggggatgcac agattcatca
 420ttttctccac cttaaaaatg cgggcattta agtntgnoca ttatctatat agnccctgct
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 539

<210> 1109<211> 373<212> DNA<213> Homo sapien
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 120ttgtactttat ggagaatatt tcaactggag gaaagactaa cttctttgag aagagagatg
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 240ctgactctca aatgaactga agatgtgccc ttacttggct gatttttttt tccatctcat
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171

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373

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201

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120ctcagtgca acactgcagc acctcagcac ctccctaacg ggggtggaag agcacttttt
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223

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180caggctgcag ctgtccaatg gcaacaggac cctcactcta ttcaatgtca caagaaatga
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326

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324

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240aatctctctc tctttttttc ttaaacaaaa tcttgggttt cgtcttaaaa atctggctta
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379

<210> 1115<211> 296<212> DNA<213> Homo sapien
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120gtgggggtgcc aacangaagc gagggacttg gtggggcggt gacttctcac tgnctccgc
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296

<210> 1116<211> 226<212> DNA<213> Homo sapien
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120gactctgggg acagggtggg acgtaccatg gggacaggag gtacacaga acacacacac
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226

<210> 1117<211> 312<212> DNA<213> Homo sapien
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60cgtttgctgt caggaaagga gaaatcactg gagagggtcca catgcctctt gggaaagcacg
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 180aggtcgggct ccatgagatg ccaatcaaat acatggggcag ccacatccct gagagccacg
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 312

<210> 1118<211> 516<212> DNA<213> Homo sapien
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 120ctgtgtccagc aattcagttt gtgctgttta agacotgggtt atttccatag agccacacat
 180gcttccaato tttcttgaag tagtgcaaat gttccttccc agagctagtt totatttgcc
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 360tttggagttg tctcccacac actcacacac acacagcctt tggaacacac natcaaaagg
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 516

<210> 1119<211> 320<212> DNA<213> Homo sapien
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 320

<210> 1120<211> 400<212> DNA<213> Homo sapien
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 120tgactctgga gctgcacagc gagggcacca ccgtcctgct cttccagttc gggatgaatg
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 400

<210> 1121<211> 337<212> DNA<213> Homo sapien
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 60ctcgtgataa agttctcttg accaaagtaa cattttatcc aaaaataaca tatctagac
 120ccgtgtagct tgacatactt caaaaatacg tgaaggtgta actgaatcac agcaggtctg
 180ttaataccga ggtactacat acttctcaaa gacataaata gtggaaagctt taagttgat
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 337

<210> 1122<211> 345<212> DNA<213> Homo sapien
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 240ctctctgcgc aacagcagca accctatcat cgtctcctgt ggctgggaca agctgtgtcaa
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 345

<210> 1123<211> 433<212> DNA<213> Homo sapien
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 60gtataaagga tatactgga actatgtgc catccctcag acttgggaag acccagggca
 120caatgatcaa catactggct gttgtgtgta caatgaccca attgatgtgt gtgaatgtg
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300agccaattat aatgatatca atgatgtcaa acggctgaaa cctggctact tagaagctac
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 433

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 224

<210> 1126<211> 549<212> DNA<213> Homo sapien
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117

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 207

<210> 1129<211> 234<212> DNA<213> Homo sapien
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 120ggccttgccg tgcactcttc ccacgctggt actttgacgt ggagaggaa ctcctgcaata
 180acttcatcta tggaggctgc cggggcaata agaacaacta ccgctctgag gagg
 234

<210> 1130<211> 347<212> DNA<213> Homo sapien
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 180ctcaggctgt aaaaagtact gagccatcct gccattcct gaggtctcta caggtgaaac
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347

<210> 1131<211> 546<212> DNA<213> Homo sapien

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546

<210> 1132<211> 169<212> DNA<213> Homo sapien

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169

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327

<210> 1134<211> 378<212> DNA<213> Homo sapien

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378

<210> 1135<211> 547<212> DNA<213> Homo sapien

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120ggaggaggagt aagttacgga gctttgaaat ttttttcatg ctttgttgat ttgaaatttg
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547

<210> 1136<211> 503<212> DNA<213> Homo sapien

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175

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96

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120catgagggc ccaagttagg tggacctgtc cctatgttaa ctgagctcgg cttaaggcc
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240ctagtactg cctagaggca catggtcccc caccagccta cagcatgaa acacccaatg
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420agtttagtcca atggacotgc ccggcggcg gcctcnaag gcgaatttca cacactggc
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527

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120catcgccctc ggcctcagtg ccatctgggg tcagaaacgt gcaggtcact ttaccccttc
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120ctcgtctaca cctctctctt ctctccact ctttatccag agtcatctcg cccttcccca
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224

<210> 1142<211> 337<212> DNA<213> Homo sapien
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337

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176

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 180cactgagaag ttatttctt gaggatagat ttccacgatg gaaaggaat gagaggtct
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177

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311

<210> 1151<211> 326<212> DNA<213> Homo sapien
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326

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159

<210> 1153<211> 357<212> DNA<213> Homo sapien
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180aacttttgcaa ggagagccaa agctaagacc cccgaaacaa gacgagctac ctaagaacag
240ctaaaaagagc acaccgctct atgtagcaaa atagtggcaa gatttatagg tagaggcgac
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357

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563

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135

<210> 1156<211> 438<212> DNA<213> Homo sapien
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438

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178

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 240tagaagagcg atottoaagg ntacatacgt gtccagctgt aagttcattt gagtagcana
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 392

<210> 1160<211> 366<212> DNA<213> Homo sapien
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 240gtcgggaaac gtgggttcaa ttgccaattg gtttctgaaa gtattcacat catttgggat
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 120aaaatncag ggg
 133

<210> 1162<211> 535<212> DNA<213> Homo sapien
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179

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 177

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180

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348

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246

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552

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120gcagtagggc ccagtagagg tggacotgtc cctatgttaa ctgagctcgg cttaaaggcc
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240ctanttactg cctataggca catggtcccc caccagccta cccatggaaa caccocaaatg
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375

<210> 1174<211> 365<212> DNA<213> Homo sapien
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180acttccacaa acaagtattt attgagcgcc tactatgtgc caggcactgt ctgacagccc
240ccccagaaa aaaaacaaaa aacaagatag aggcagcaaaa cacaattctc gagggagagg
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365

<210> 1175<211> 583<212> DNA<213> Homo sapien
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583

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120gggacacact ttgggcagct gtgtgttcgg gcattggacg ctggtcagtg ttccgtcct
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 420atgctgtgac gctcgacagc ttactaccan caggggcccc caaggangga gacaaacctt
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<210> 1178<211> 395<212> DNA<213> Homo sapien
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 240tgaaaaatat ctggtttatat cattctgggt gtgtttctac tgacacagg ggtcccgctg
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<210> 1179<211> 196<212> DNA<213> Homo sapien
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 196

<210> 1180<211> 635<212> DNA<213> Homo sapien
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 180acatctgcaa agactgttac gcaacacag tgcgtctcgg cggcaccaacc atgtaacctg
 240gcattgcga caggatgcag aaggagatca ctgcctggc acccagcaca atgaagatca
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<210> 1182<211> 694<212> DNA<213> Homo sapien

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 556

<210> 1184<211> 363<212> DNA<213> Homo sapien
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 120ccaaagcaac agtgttccaa cggattcaaa ggggtggcatt gggttggagc tctctgggtac
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 363

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<210> 1186<211> 499<212> DNA<213> Homo sapien
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 499

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180ctggtgtctga gtcgcgcagt cctcgccatc ggccaccga tctcagcatg gcacagcaga
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 290

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 570

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 234

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 120ggtcattcca gaaggccagc gagaaactgt gggaaataat aaaacctccc tctccacat
 180ggcgcgcaagt gctgttttaa gcaaaactct catttcaatg tgagggttag aaaaactatt
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 279

<210> 1193<211> 335<212> DNA<213> Homo sapien
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120gaagactct catgaattaa atagtgtgat caatttttaa cgttaattga tataaaaaaa
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 240ccagacatac tgtgttgaga gatacttga gggaggaggt aggttttgaa gaggttgatg
 300gtgtgtggga ggaagagacc tcggcccgcc accac
 335

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 306

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 372

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 612

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 284

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 240ttagggatcan caaantaaag ccactaggc tgaggatcac cctcactcc tttactccat
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 347

<210> 1199<211> 190<212> DNA<213> Homo sapien
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 190

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 360agg
 363

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 420tatgtgtgta taacatttac ctgcccnggc gggcgctgac aaagggcgaa ttccaacaca
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<210> 1205<211> 103<212> DNA<213> Homo sapien
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103

<210> 1206<211> 458<212> DNA<213> Homo sapien
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<210> 1207<211> 431<212> DNA<213> Homo sapien
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<210> 1208<211> 747<212> DNA<213> Homo sapien
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<210> 1209<211> 213<212> DNA<213> Homo sapien
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 180cgcaaatacc tgcagggcaa acggggcatt gag
 213

<210> 1210<211> 743<212> DNA<213> Homo sapien
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 280

<210> 1213<211> 342<212> DNA<213> Homo sapien
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 342

<210> 1214<211> 294<212> DNA<213> Homo sapien
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<210> 1215<211> 371<212> DNA<213> Homo sapien
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<210> 1216<211> 654<212> DNA<213> Homo sapien
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360cttggaaagcc ttgaatgtcn ttaaccggaa taaaagggtc ccattgcttc caaccccga
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479

<210> 1218<211> 173<212> DNA<213> Homo sapien
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173

<210> 1219<211> 201<212> DNA<213> Homo sapien
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201

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506

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248

<210> 1222<211> 381<212> DNA<213> Homo sapien
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381

<210> 1223<211> 446<212> DNA<213> Homo sapien
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446

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240

<210> 1225<211> 246<212> DNA<213> Homo sapien

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319

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268

<210> 1228<211> 618<212> DNA<213> Homo sapien

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618

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267

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291

<210> 1231<211> 326<212> DNA<213> Homo sapien

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190

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326

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256

<210> 1233<211> 312<212> DNA<213> Homo sapien
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312

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331

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380

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372

<210> 1237<211> 102<212> DNA<213> Homo sapien
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102

<210> 1238<211> 467<212> DNA<213> Homo sapien
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120cgtcagctc taaaggttac tganogttaa tggagggcgg gagcangaag aaagtcaang
180acctggcaaa aagatcatct tccctccata tctcttctga ggtaatatta ngtaaacang
240nacctggacc agagggctca attatataca tagtaacctt tattctgaat taacattata
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360tggcgaagtq ggagggagga catnatgtta ggagccctgt ttgggggaagg aaatgttttc
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467

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120atgcagcttt tgcataaagcg ggggcggct tcctctctag ccctcagct tgcctaccct
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264

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176

<210> 1241<211> 301<212> DNA<213> Homo sapien
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180tggctagcgt gagagcagg atctacctgg cttctcagtt ctttggttgg aaggagcagg
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300g
301

<210> 1242<211> 108<212> DNA<213> Homo sapien
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108

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142

<210> 1244<211> 559<212> DNA<213> Homo sapien
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180ctctccagatt tgcagaagaa aatccacatt ttgcccaaa atgtctaat cttgacggag
240agtctaaagc agtccacaga aaaaatgcag tcagatatgg agaaaatcca agaatcaga
300gaggtccagt tatactcagt ggaagtgact ctggaccacg acacggccta cccagcctg
360atctctctgt ataactgcg gcaaggtgcg gtacagttac cttaacagg aactgcctga
420caaccocagg aggttcaatc tgtttccctg tgtcttgggg ctctccattg ctctatgcc
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559

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120tgttttaatt gccccaacca tngaaacaaa atagaacctt aaataaaggt caggggttaa
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277

<210> 1246<211> 256<212> DNA<213> Homo sapien
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180ccggcagttct tggcatcaac cacaaagcct acttctctgc cagttttcac agtggaggcg
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256

<210> 1247<211> 550<212> DNA<213> Homo sapien
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550

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108

<210> 1249<211> 240<212> DNA<213> Homo sapien
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120ttactaaaaa gccttcactg gtttggatgc attgagact ctagacctga tgcacaacgc
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240

<210> 1250<211> 553<212> DNA<213> Homo sapien
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120accocgtctc octaaaaata caaaattagt tgggcgttgt ggcgcagtcc tghtaatccca
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553

<210> 1251<211> 246<212> DNA<213> Homo sapien
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120gtactgcagc agcgaacatc tcgatatatg aaaaotgcat catcaattca acgttttgggt
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246

<210> 1252<211> 550<212> DNA<213> Homo sapien
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550

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<210> 1253<211> 245<212> DNA<213> Homo sapien
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 120gggggatgcaa acgtgcaaaa gcaggggggaa gctgcccagc ctgagactgg agcagotagg
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 240gcagg
 245

<210> 1254<211> 556<212> DNA<213> Homo sapien
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 120gcattggaggt cctgtggcact ccacgaaact acctcaact ccactatgaa gtgtgacgtg
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 240ggcattggcc acaggatgca gaaggagatc actgcccctgg caccacgac aatgaagatc
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 480ttgcaaaacc taacttgenc aaaaaacaan atnaaatagg catggctttn tncgttnttt
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 556

<210> 1255<211> 494<212> DNA<213> Homo sapien
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 494

<210> 1256<211> 312<212> DNA<213> Homo sapien
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 180ttctggctg cantcncan ttccctgcon tngggcacc gatctacna nggcacagca
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 312

<210> 1257<211> 441<212> DNA<213> Homo sapien
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 441

<210> 1258<211> 287<212> DNA<213> Homo sapien
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 287

<210> 1259<211> 339<212> DNA<213> Homo sapien

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 339

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65

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 177

<210> 1263<211> 560<212> DNA<213> Homo sapien
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420aacccogaaty gacanggtaa anaangaaty ggaanaggca gantttcaag ctaagaactc
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549

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246

<210> 1267<211> 143<212> DNA<213> Homo sapien
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143

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447

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223

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555

<210> 1271<211> 452<212> DNA<213> Homo sapien
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452

<210> 1272<211> 549<212> DNA<213> Homo sapien

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196

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 240caagacaga gaaaataccc anagttngga gacaggggga acatctcanc tacaacact
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 120tccaggtcgg agtgcaatgg catgatctcg gctcaactga accccatct cccaggttca
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 240ctaatttttt gtatttttag tagagacggg gtttccatt gttgaccag ctggtctcga
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 330

<210> 1274<211> 535<212> DNA<213> Homo sapien
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 120agctcctcag cagcctgact gactgcctga cggtaggccc cctcagtgcc agcgtctgga
 180ggcagctgta cctcaagcac ctgtcacagt ccagccttct gctggagcac ttgctcagct
 240nctgggagca gattcccaa aggttacaga agtctttctc agaaaccatt cgtctcctca
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 360gtgacatgg ctgcaagggc ctgttcanc aggttcang tctctcgmt cccctggagc
 420cggctnctcc tgtggtcgt ggnctctcg nanngctcc ttggtccatg acctcccnct
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 535

<210> 1275<211> 262<212> DNA<213> Homo sapien
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 120cagatattgt ttggagtgc ttcttcgata ctacagttat ggcttggaaa aagaattccg
 180gtggagacata tcaaggatt tccaggagga aacgtggaag gactatgaag ctggtaaag
 240ccagagtgg atctgagtga gg
 262

<210> 1276<211> 289<212> DNA<213> Homo sapien
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 60gggaatgact tgagtttgag tacagatatt cgacaccaa aaagtcatc tacaatgaat
 120tcttatgaat gttatcaatg tgggaaagcc ttctgcgaaa gttcatcctt tattcgact
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 289

<210> 1277<211> 90<212> DNA<213> Homo sapien
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90

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 120acatctgtat tttttatcta tatctgccct gatgtcagtt atagattcca cagtcttgat
 180ttctttggag ctgcattgg ggaaaaggtg tgtgtctcag aagttataaa caaacctcat
 240gagctggagc ctgtaacgtg ttgcatctct cgtgaaatgt ggagngagtg tatgtcctct
 300tccaaaagca atcacagat tggggctcna agtgttctct ttccacang agctggggtt
 360gagcaccaat gtggaatntg atggcaggtc aaaggtcatg ttcttatgct cactcttggt
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480agggcnaatt ccacacactt tcggcccg
509

<210> 1279<211> 381<212> DNA<213> Homo sapien
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120tcaaaagatt ggtataataa caaatgtttt caaatgtatt cgtctatac aagacagatg
180ttttatgtga cttgtgcacat aactatata catcacataa taatatatta tgaattattt
240cccaagatt gagcgacac atagggagaa aatytaaagt tctcaatttt gtgtccaaa
300atgattattt atcaaaattgc tatagcgtgt ggatagctta aagaaaaaaa agtttctcta
360aatctggga caaacagatt t
381

<210> 1280<211> 120<212> DNA<213> Homo sapien
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<210> 1281<211> 327<212> DNAC<213> Homo sapien
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 60taactcctgagc gttgtccgtc taatgacata ctgcgatcc tgaatactgg ggtcattttg
 120ttttttcttc tcccaagttt agatagcgg aacgcgcgaa ggaagcagc catctggcgt
 180taatttcatt ttaagttttt ctgttgataa aaatgtgaa ttccatagcc catctggcgt
 240aaggtttctc gtaactgttg ctgtctatgt aaatgactcc ctcaaaaactc agttttatca
 300gcacgtgcgc tctgtgtgag tacatgg
 327

<210> 1282<211> 432<212> DNA<213> Homo sapien
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 120gtgaaaaaatgt ctttttccct cctctcagac accgttttat atatngcga gaataatgaa
 60ttgaatttgtgt gtgtcctatgt ggggcnnaat tnaaaattac caacgaagc cagctgttgt
 180atcnacccctc cctctcctct aggaactctaa aaaaaaagaa aaaaacnctt cagctgttgt
 420gtgcaaaaaat ttgaggaacc cctcttcacac agtacagcat ttaatngtc agtgcamaa
 300aatctacatct tgcncacaan tgttttgaca tgatttcttt tacnaaacct tnaagtcmaa
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 180agg
 183

<210> 1284<211> 261<212> DNA<213> Homo sapien
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60cattctctgac aagagatcga gccctcttgc tggcctctgc cactatggtg aagtgaacaa
120tgacttttgc atgctgaaag ctggtgtgtg gggaaacaa aagcgggtgc tcaccttcg
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261

<210> 1285<211> 222<212> DNA<213> Homo sapien
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60aaataatccac cagaggggaaa tctctgtgga agggatgaa cggcgatttt gctacacctg
120ttatgtatgc aaagaaggag acctctctct tgaagaggaa tctctgtcac tctctcatgc
180tgtctatgct atacaacctat ttctcttttc ttacaggatt tt
222

<210> 1286<211> 479<212> DNA<213> Homo sapien
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60aatgtgggtc atctctcttt agcaattttg cttagaagaq actgaattgq qtttttattt

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198

120gctttttgat atocaaataa tgttttttcat ttttttattc otttataagt ggaaataaca
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 240taacgaacaa tctctatgct tctttacact tttaggaaga gaatacactc caattctgca
 300agtgtgtcta tttctagag gtytaggtgt gctaagaatg tgattccang actgcotttt
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 479

<210> 1287<211> 310<212> DNA<213> Homo sapien
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 120tgtctctggg gctcagccccc agatggggcc agagcaggga catgaggacc gaatggtgcc
 180gctcattccac agcattgaga ggcctggct caggagtccc cttctccagc tctctctgga
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 310

<210> 1288<211> 528<212> DNA<213> Homo sapien
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 420gagataaen totgctatca agagggttca taaatttatg gaacctgtga aggtccagac
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 180ttgtactgaa ttctgtcag ttaagggggt caactgcttg gtgaaaattg gtgaaaattg
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 383

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 120gttgagccct agccatcact gtctcttaac ataatttgc atcaaaaaga caaagcaaat
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 240gctttttcat tabaaaaat aaagacatta aaatgcgaaa acaaatgacc tggagggaaga
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<210> 1291<211> 377<212> DNA<213> Homo sapien
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 240ggatcgtgat ggcattggga agctggccct ggtggagttc aacatcctgt ggaacccgat
 300ccggaattac ctgtccatct tccggaagtt ttgctggac agtcgggca gcatgagtgc
 360ctacagagtg cggatg
 377

<210> 1292<211> 473<212> DNA<213> Homo sapien
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 60agtggtgggt cctgggtgga aaccagcttc cacatacac ccccaaggga agaattgggg

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199

120ctctttctgga atggaagaag ctgacttcaa gcatcttcaa aattactctc gttcccacgc
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240nncaatgggg ataangcata tggctgaggac aggggagagc gggaggggca toatnctgc
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360gggttgagga gtnagactga ggcanataaa tacccttccc catccttaac tctagaaccc
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473

<210> 1293<211> 536<212> DNA<213> Homo sapien
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536

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120caaatggcctc attggagatg cgtcagtgca tgacgtgtct cttgtctggtg ttgcgatga
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425

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325

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314

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204

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200

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 539

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 512

<210> 1300<211> 549<212> DNA<213> Homo sapien
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 420gaaagaagc agaggaat ttgatcagntt gttagccgag gaaaaaaca tctcttncaa
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 549

<210> 1301<211> 532<212> DNA<213> Homo sapien
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 532

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 273

<210> 1303<211> 281<212> DNA<213> Homo sapien
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 120aacacttagta tgcagttgat aagaggaatt tggatataat atggtgggtg attatttttt
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 281

<210> 1304<211> 315<212> DNA<213> Homo sapien
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201

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 240agttaactga ggacaaggca gacgtgcagt coactatcgg cctgcagcgc ttctctcgaga
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<210> 1305<211> 180<212> DNA<213> Homo sapien
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 60gtcaactaata gaatgtctcc aactcggat tgaatggag aaaaacacct tccctctag
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 180

<210> 1306<211> 184<212> DNA<213> Homo sapien
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 120gtaccocgag atgagactgt gaggtacggg ccccatcaca tgggtctaac ataactcgcg
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 184

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202

305

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203

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384

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112

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287

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325

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308

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242

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382

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546

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 328

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 378

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206

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 317

<210> 1340<211> 543<212> DNA<213> Homo sapien
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 360acaaatgtat ttgcaaatgt tcacaaatgc taacaggagg ctggtgtgtt gatcagatg
 420gnccttccaa cttgaacgga atgtactatc cacagaggga gaacacaaat aagttcaacc
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 540cca
 543

<210> 1341<211> 536<212> DNA<213> Homo sapien
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 240cagtcagcat ctattcttgg gcaccgggag gagcgctct ctggaatctt aaattaccag
 300gtaggagtgga gaatcgtgat tagctgtaga gctgaaggca cggctgcag gccctcagg
 360gtcctctctc anttctccag gcccggtgta tcgaangcag tggaaaggaa gggcctgtgg
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536

<210> 1342<211> 539<212> DNA<213> Homo sapien
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120aaacttcga agcccgttct tataagtgtc tcatctctta cctgggtgta aatggaatgt
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240gtggactct ctttgactta aagggcgat gatgaataa aactcaaacg cctttctctc
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360ccgtccaaag cggccatggc ccattgtttc actagatggc gtgcacatt caggcatcaa
420ccctcatggc ctctcagoot tgcaaaaggca gccacttaaa gtgcgtgtcc tgtgtggggc
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539

<210> 1343<211> 224<212> DNA<213> Homo sapien
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120ctgcgtttggc ggtgcagtat tcttcatagt tgaacatata, gctggagtgg tcttcagaat
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224

<210> 1344<211> 408<212> DNA<213> Homo sapien
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120aaggtgcgat gatgaactac agtcaatgac ttagacaaag gcgatgccag tggggcttgg
180tatgttctca agcatcaatta cccatgccat cccattccag aggttgtgga cagcgtctgt
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408

<210> 1345<211> 177<212> DNA<213> Homo sapien
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177

<210> 1346<211> 219<212> DNA<213> Homo sapien
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219

<210> 1347<211> 538<212> DNA<213> Homo sapien
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240ataaaaaaaa agtagtcaata agacaaaata acaaaaatta tctottaaga gttaaaaaat
300aggttgaaag tggttgacca ggtgatatto aagtttagtt tagtcgaaag cttttacc
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538

<210> 1348<211> 290<212> DNA<213> Homo sapien
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240gaaactggga tgaacagttt gacaaggaga acaccggaga gagactgttt
290

<210> 1349<211> 540<212> DNA<213> Homo sapien
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120cttgaggagct tataaaaaa agtcatctta agttcacaa ttgtacaaga agaaagtgtt
180ggataaactag gaattatttg taagtaaatgt ttcaagttag tactagcaa tatagattct
240tttattaaga tgtatctgtc gaattaaggg taacagttga aatagtctg tggcgtctct
300aagaataaat gggaaaaaaa tctctggatg taagtttttc tgtgaaact agaggggttt
360ttttttctgt ttncatatac ttttttttaa tagcaatggg tttttattaa acagtgtggg
420ggccacaggg catgttgttg gngaaatata taaacattta ttacctcgg ccgncaccac
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540

<210> 1350<211> 243<212> DNA<213> Homo sapien
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120cacttggcac acaggtttgt atgtatgtgt atatatatgt gtatgtatgt atgtgtggg
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240att
243

<210> 1351<211> 382<212> DNA<213> Homo sapien
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382

<210> 1352<211> 535<212> DNA<213> Homo sapien
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120tgacatccca gatgagggga ggctgttcc agaaaacccbt tcgcaccogg tctttatcac
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360gatgaccacc atentacttt ttgtatgggc attctccca ntgcactgt ttccogman
420cccggtgcc cgttccccn cgaacccacc caacccggcg gncnacottg gaaggcttc
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535

<210> 1353<211> 412<212> DNA<213> Homo sapien
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240agtacagaat ctgttctatt ctgctccag aaaatcgaaa acctgtgagt cagagtca
300gaactttacc caagcaacgt aattcctgt ttcatgggtc ctgtagatgt ttgattcan
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412

<210> 1354<211> 85<212> DNA<213> Homo sapien
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85

<210> 1355<211> 427<212> DNA<213> Homo sapien
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120gcaaggggga ggaaggtac tgaatacac tttatgaagc aagtgtgtct cgggctgtgc
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<210> 1356<211> 266<212> DNA<213> Homo sapien
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 180tggtgattag tcggttgttg atgagatatt tggaggtggg gatcaataga gggggaata
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 266

<210> 1357<211> 343<212> DNA<213> Homo sapien
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 240ctgtgggtt tttctgttaa nactgagga tgacttntat gattggcgcc ancaagtcaa
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 343

<210> 1358<211> 102<212> DNA<213> Homo sapien
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102

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 486

<210> 1360<211> 181<212> DNA<213> Homo sapien
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 180g
 181

<210> 1361<211> 269<212> DNA<213> Homo sapien
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 120ctaagtgccc aacatgaaca aattaaaacc ttaataaagg gtcactgtta acgcctatcc
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 269

<210> 1362<211> 124<212> DNA<213> Homo sapien
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 124

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210

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276

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270

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180

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211

<210> 1367<211> 179<212> DNA<213> Homo sapien
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179

<210> 1368<211> 384<212> DNA<213> Homo sapien
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240ccagattgtc ccactgttca cagatctttt gccaacgggc gttgacctg ggtgagtcac
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384

<210> 1369<211> 241<212> DNA<213> Homo sapien
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240
241

<210> 1370<211> 302<212> DNA<213> Homo sapien
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302

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211

<210> 1371<211> 277<212> DNA<213> Homo sapien
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 277

<210> 1372<211> 462<212> DNA<213> Homo sapien
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<210> 1373<211> 241<212> DNA<213> Homo sapien
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 240g
 241

<210> 1374<211> 133<212> DNA<213> Homo sapien
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 133

<210> 1375<211> 495<212> DNA<213> Homo sapien
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 240ttttattaaag tgcattctgt ggaattaagg tcaaggttga aatagtctg tggctgcta
 300agaataatg ggaaaagaat cctcggatgt aagttttctt ttgaaactga gagggtnttt
 360ttttctgttt acataactt tttttaatag caatgggntt tttattaaaa catgctgngg
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 495

<210> 1376<211> 110<212> DNA<213> Homo sapien
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110

<210> 1377<211> 171<212> DNA<213> Homo sapien
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 60ccaaggacac accgtagtct tccaggaaga tcactttcaa cctgtcacc acaactgggt
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 171

<210> 1378<211> 494<212> DNA<213> Homo sapien
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494

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406

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300aacaacagcgt ctgttgcaat gtaacttgtg gctgtgcatt aagatgttgc ntgaggattg
360ccaactcctg caccatattt atactgntgg aacggtgcgg acagcaggag taacttgcac
420cgctncaag ctncaggacg tgggacccat ttgntctgtg ttgatgtggn naanaacacc
480cttngtnnga cttnacttct gggaaccnn
509

<210> 1381<211> 256<212> DNA<213> Homo sapien
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120cagtgaggag gatgcocaaag ctgccaanga ggccatggaa gacggtgaaa ttgatggaaa
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120catctgggtt cctcaaggac agggaaactgg ggaactggct caaagttagc atagaaata
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300ncttnatcct tgnataccaa ggaagctnt tacttgcnt ggocctcaa gaactctggg
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441

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120catctcgaat ctgtccctt ttggggtaact tgggggagac tctggctccc aggatctgtt
180cctatttca gtgccttctc angacacagg ggaactcctn acgtcccaca ggtttctnt
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296

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180aaggcagttaa caagaagata gatccacaaa agacattaca acaatgcag aatttccaga
240aaggaaacat gaaaattgaa atgactgaag aatatgcaa tgatacact gtgacatct
300ttgcaggttc tgatgacgaa gaagaaagcc aggatattgt gaatcaagtt cttgatgaaa
360ttggaatttg aaatttctgg aaanatggac ctggcgcgcg gaccac

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213

406

<210> 1385<211> 504<212> DNA<213> Homo sapien
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360aacctttctct ttgccatttc ttctttctct tttttaactg aaagctgaat ccttcatttt
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504

<210> 1386<211> 488<212> DNA<213> Homo sapien
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120gggtgtggcg cgaagatatg ctcatgtggt gttgaggaaa gcagacattg acotcaccaa
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488

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180tatggagtgg atcactgcaa aactcaagga ggcocggggc agaggaaaaa aatttaccac
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502

<210> 1388<211> 508<212> DNA<213> Homo sapien
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240atacccaaaa gaaataaata tatacccaaa ggaatgaaa tatttgcaa tatncaaaa
300gtagaagtaa cccaagtgtc cattgtctga ggggatggat aaccaagat gtggnacata
360catatttaag aagtattatc cnccttaaaa ggaatggaa ttttgaccoc taactcact
420tgggatgaac cnccaaaaan attattatta ttattggnnt tatttttttt ttttttgaa
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508

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120aaggcctctnc aaagagatgg accgaacctt gggttaggca gccctctgc ccaagagaaa
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360cncantntga tnaoctggga agaattacca ctctccctgc ttaannnacc aagcagaaga
420gtcttttttg naagggggcg ggnattgggg ncnattattt ncccccnctg gnnttttctc
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539

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214

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326

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120ggcctctgcg tgcctccttc ccacgctggt actttgacgt ggagaggaa cctgtccaata
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234

<210> 1392<211> 403<212> DNA<213> Homo sapien
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300tcagtataac ccggggaggg accttccctg cctctgtcgg ggtgctcttt ggacactgga
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403

<210> 1393<211> 504<212> DNA<213> Homo sapien
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504

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120ctaagaaagt ggcctggaga tgttttagaa gtttaaacca acgaagaaga aaatcaatga
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267

<210> 1395<211> 378<212> DNA<213> Homo sapien
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378

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120tcaaaagcca actctgagga gacgaagtgg caaagaacag ccttggctc cctcccca
180agaagaacgg cagctgcagc tgcctggaaag ggcaagaate agagtggggg gaacactcgg
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215

259

<210> 1397<211> 508<212> DNA<213> Homo sapien
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<210> 1398<211> 409<212> DNA<213> Homo sapien
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 409

<210> 1399<211> 199<212> DNA<213> Homo sapien
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 199

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 439

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 570

<210> 1402<211> 294<212> DNA<213> Homo sapien
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 120attttgaaa aggcactca cagttgcttg tgggttatga aagaattggc cctacgtctt
 180gcattgaaa tgttacaggg gacattgggc caggcaattat tatatagaga agtctattt
 240ccaagctct gactaacttc tggatatgaa aataaggaa cttgccagca tagg
 294

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216

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103

<210> 1406<211> 384<212> DNA<213> Homo sapien
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 384

<210> 1407<211> 226<212> DNA<213> Homo sapien
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 120caactgttgc caaagagttg gctttgttta ttgtgtttt gggggagag gagtggatt
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 226

<210> 1408<211> 413<212> DNA<213> Homo sapien
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 360cccgcgga cggagggcgt atggtgaact gacnagcttc anacaccagct ggg
 413

<210> 1409<211> 441<212> DNA<213> Homo sapien
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217

120gccttctgtg acagcaccag ctggacgttg tcatgaaatg tcaogagttc tgggtgtttc
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 441

<210> 1410<211> 453<212> DNA<213> Homo sapien
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218

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218

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122

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686

<210> 1421<211> 569<212> DNA<213> Homo sapien
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 413

<210> 1423<211> 643<212> DNA<213> Homo sapien
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 120cttcgaggca accacgagac agacaacatg aacagatct acggttctga ggtgtaggtg
 180aaggccaaagt acacagccca gatgtacag ctctttagcg aggtgttoga gtggctcccg
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 284

<210> 1425<211> 243<212> DNA<213> Homo sapien
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 243

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<210> 1428<211> 691<212> DNA<213> Homo sapien
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 60tgggtgccatg acaatggtgt gaactacaag attggagaga agtgggacgg tcaggagaa
 120aatgg
 125

<210> 1430<211> 116<212> DNA<213> Homo sapien
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116

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 133

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<210> 1434<211> 294<212> DNA<213> Homo sapien
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 294

<210> 1435<211> 674<212> DNA<213> Homo sapien
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222

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222

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725

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294

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240ccagatttgc ccactggcca cagatctttt ggcaacgggc gttgacatg ggtgagtc
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390

<210> 1444<211> 156<212> DNA<213> Homo sapien
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120aaantccagg ngncctctct cncnctcnc cctggg
156

<210> 1445<211> 706<212> DNA<213> Homo sapien
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240tccaactacat tgatgagcta ctcatccag actcagccaa gacactattt gaattggctg
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420ccccctcaat tgatgcccat acaaaggaatt tggctggaaa ccacataatt aaagaccaag
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706

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223

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 304

<210> 1448<211> 637<212> DNA<213> Homo sapien
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 360gtgtgaaact gcatctctac tggacatgag ggccttccat gtaggacaag aggtgagttc
 420gtttattttt gtaactgggt tacaagtctc gattagttaa tcnngagctt attgtcatct
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 637

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 180ggcgccaaag cgtgttttaa gcaaaatcct catttcaatg tgagggttaag aaaactatct
 240tgggttcagg gtatcctttg tccaaggtac gagaggagg
 279

<210> 1450<211> 317<212> DNA<213> Homo sapien
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 120gaaagtctct catgaattaa atagttgatg caatttttaa cgttaattga tatataaaaa
 180aacacaaaaa ttaggcttgt aaaaactgact tttctattac gtgggttttg aaactancc
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 317

<210> 1451<211> 297<212> DNA<213> Homo sapien
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 120tgactctttt ggggaaccaog tcaaccaagt acacaggga gaaagccatg tatttacat
 180ggcagggttc acatttcacc atctggttgg ctggctcaaa gcaagcatlg gtatgctctg
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 297

<210> 1452<211> 445<212> DNA<213> Homo sapien
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 120aaattctttt tctataacag ttccagaaaa agcctcaggt gttactgata agggcaaaag
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224

300gctttcacta gtgtogctac acacactatt ctctctaact cgagacttca ggtatggatt
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445

<210> 1453<211> 302<212> DNA<213> Homo sapien
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300tt
302

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120gtgcagtata aaatataaaa aggtttgatt ctgaatagac caactgtcaa ttttctttaa
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240taagagtctt atttgcaata caaaaactgg agcttatgac tgctttgatt ttctctgtag
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372

<210> 1455<211> 310<212> DNA<213> Homo sapien
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310

<210> 1456<211> 344<212> DNA<213> Homo sapien
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120cagtttagtc attaaagttt tggaaattct cagacagatgc agtggatata gaaacttgta
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344

<210> 1457<211> 332<212> DNA<213> Homo sapien
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332

<210> 1458<211> 540<212> DNA<213> Homo sapien
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540

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225

<210> 1459<211> 223<212> DNA<213> Homo sapien
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120cttgcccttca tttactcttag tccaagattc ttgcaaaaca ggaactgaa caaacattag
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223

<210> 1460<211> 368<212> DNA<213> Homo sapien
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60gttccaaatca cgtatgagtt tttctaaaaa aaccgaaaca ctggaacat ggatgcactt
120taagaacttt atgctaagtg aaaccagtca caaaaggaca aataactgtat gattccaactt
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240ggaaggggag gtggggagtt attgttcaat gggcacagaa ttggggaaga tggaaaactt
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368

<210> 1461<211> 290<212> DNA<213> Homo sapien
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120agctaactgc caggaaagaa aatttcoctca taaatactaa gcaacttttt cattacaactg
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290

<210> 1462<211> 535<212> DNA<213> Homo sapien
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535

<210> 1463<211> 484<212> DNA<213> Homo sapien
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484

<210> 1464<211> 267<212> DNA<213> Homo sapien
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267

<210> 1465<211> 231<212> DNA<213> Homo sapien
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60ctgtagtgtt agggcataga actgtgaggt cagatagtcg gccacgggg cgggtctgt
120gacctgtgac gccacagggt ctctctggcg cagccttgca aatcccacca cacaggtctt

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226

180ctctctcaga tgcagaagca ctttggaact gcccgcgccg cgctogaagg g
231

<210> 1466<211> 202<212> DNA<213> Homo sapien
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120tggaaactgt tttttttctg ctttgtttt toagtttctg gttctgtag ccatattgta
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202

<210> 1467<211> 97<212> DNA<213> Homo sapien
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97

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342

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308

<210> 1470<211> 284<212> DNA<213> Homo sapien
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284

<210> 1471<211> 490<212> DNA<213> Homo sapien
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490

<210> 1472<211> 286<212> DNA<213> Homo sapien
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180catgccctc cctccagaa aggcagaaaa tggagagaat gctgttaagt aagaacccaa
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286

<210> 1473<211> 230<212> DNA<213> Homo sapien
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227

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230

<210> 1474<211> 330<212> DNA<213> Homo sapien
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330

<210> 1475<211> 197<212> DNA<213> Homo sapien
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197

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326

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538

<210> 1478<211> 288<212> DNA<213> Homo sapien
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120cttagaagtg aggtctgtgag caggagcctc tgcagggga tgcacatct gtggggaggg
180gccgaggag actccatggt ctctctgtgc tgcctgtccc tctctgtg agaaagagctt
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288

<210> 1479<211> 141<212> DNA<213> Homo sapien
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141

<210> 1480<211> 388<212> DNA<213> Homo sapien
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120caggaggttgc agtgagccga gatcacgcca ctgactcca gcttggcgca cagagcgga
180ctcatttca aaaaaagaa ctacaagttc tgattccgga ctccagatg tgaattttaa
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228

300atgtctggagt caaatctgaa ccccaacta tgcctctta aggggggtcc ctctgggatg
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398

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540g
541

<210> 1482<211> 424<212> DNA<213> Homo sapien
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120aaatgaaggt gtctagtagt ttgttatagca cggtagtcca ttctttctaa aggaccactt
180acggtttcatg caggtaatat taggtatagc ccatcttcca ttctctaagg accggaatgt
240tggaaggtga cgttgaatga aaccatcttt gcaagtgttat ctaatcaggg agttgatttc
300ataacagaggt ttcatctttc caaaggcttt ggcatttttt caacacaggg cctggccgca
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420ctgg
424

<210> 1493<211> 431<212> DNA<213> Homo sapien
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180ggttttccag gggagggggt ttgtctgtaa gggcaggttc agatgcagcc ttccagattt
240aaggagcact ggaggacagt ggcctgagtg aggcgccac acctggccc gcacaggct
300caggcccaca ccttgtagt ttgaaacca aagcccaana gatgatgttt acttctctct
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431

<210> 1484<211> 99<212> DNA<213> Homo sapien
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60gaacctgcag ggaagtactc cggaaactgg gggtagcagg

99

<210> 1485<211> 192<212> DNA<213> Homo sapien
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192

<210> 1486<211> 98<212> DNA<213> Homo sapien
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98

<210> 1487<211> 255<212> DNA<213> Homo sapien
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120ggagttgttt tcagctataa caggattacc cgcagagct gtgctacaaa cagacaccag
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255

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<210> 1488<211> 261<212> DNA<213> Homo sapien
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 120cgtgaacttt caagtgtatg catctacta ctgagaagtg agagagaggt cttaaaggggt
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 261

<210> 1489<211> 344<212> DNA<213> Homo sapien
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 120cagtttagttc attaaagttt tggaaattct cagacagtgc agtggtatca gaaacttgta
 180ttcaagagta caggtcagag tctctctttt ttctcttttt gagatggagt cttgctctgt
 240tcacagactg gagtgcagtg gtgcgatctg ggtcactgc aatctccacc tcccggttcc
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 344

<210> 1490<211> 426<212> DNA<213> Homo sapien
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 180cgttgatga ctgccttttg tctgggcttt ggagggccct tcccagatgt ccagcctctg
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 300cagcgtcaat agagtgcagc gcatggcgct cctctccacc actgcgtgc ggaaccaaac
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 420gaagta
 426

<210> 1491<211> 339<212> DNA<213> Homo sapien
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 339

<210> 1492<211> 543<212> DNA<213> Homo sapien
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 120gttgttggtg atgaagggtt tgggtggctc atagactgtg atcgtctgta ctgtgtctct
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 300gtactgtgca ggtgggttag aggtcgtggt caggaagaggt ttcagatttt cccctgatct
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 420gtactgaatc actgcctcgg gactcactgg gttctgggt ttacattttg tancttgctn
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 540cct
 543

<210> 1493<211> 77<212> DNA<213> Homo sapien
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77

<210> 1494<211> 344<212> DNA<213> Homo sapien
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 60tatgagtgtg gaatccagaa caaattaagt gtgacaaca gcgaccaggt catctggaat
 120gtctctctatg gcccaagca ccccaactt tccctcatt acactatata cctgcagggt
 180gtgaacctca gctctctgt ccatgcagcc tctaacacc ctgcacagta tcttggtgtg
 240attgtatggga acatccagca acacacaca gagctcttta tctccaat cactgagaag
 300aacagcggac tctatacctg ccaggccaat aactcagca gtgg
 344

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<210> 1495<211> 380<212> DNA<213> Homo sapien
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 120gcactccagt ttggcaaacg agcaagactc catctcaaaa agaaaagaaa agaagactct
 180gaactgtact cttgaataca agttttotgat accoactgac tgcctgagaa ttccaaaaac
 240tttaataaac taactgacag cttoatgaaa ctgtcoacca agatcaagca gagaataata
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 380

<210> 1496<211> 540<212> DNA<213> Homo sapien
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 180cgcttggcag gatgatgtgt toattagat ttcaacaaga gtacttccag agggtaactt
 240aacagagatc cagatctatc ttgtcaatcc caacgtttta cataaaataa gagatctctt
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 420accacagcat gcaatgccaa ataataaat tgcctcacc agactccggg ncogagaacc
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 540

<210> 1497<211> 212<212> DNA<213> Homo sapien
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 120tgccttgata gtccagtcaat tatttgtgta tgaacaatg tacaaatcaa ttttttgaaa
 180taaatgatct cagaactttc aagttaaatt tt
 212

<210> 1498<211> 204<212> DNA<213> Homo sapien
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 60ggagcccaga agcatccagg tacactttcc aaacaggcga cctaccagg aactgggaga
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 204

<210> 1499<211> 305<212> DNA<213> Homo sapien
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 120agcaacacgt agacattcat acatatccgg tgaagaagct gtttctgaga tgcgattgcc
 180atccaaacgc aaatgcttga tcttggagta ggataatgcc ccaggatctc tgcagaagct
 240ctttatgtca aacttctcag gttgattgac ctccaggtaa tagttttcaa ggttttcatt
 300gacag
 305

<210> 1500<211> 547<212> DNA<213> Homo sapien
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 547

<210> 1501<211> 53<212> DNA<213> Homo sapien
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53

<210> 1510<211> 415<212> DNA<213> Homo sapien
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 60gttggttctt gagggtctgc caatgacaac aggacctcca ctctactcag ngtcacaagg
 120aatgctatgag gacctctatg gttggaatc cagaacaaat taagtgttga ccacagcaac

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232

180ccagtcacac tgaatgtcct ctatggccca gacgacccca ccatttcccc ctcataccac
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300cagtatctct ggctgattga tgggaacatc ctgcacacaa cacaagagct ctttatctcc
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415

<210> 1511<211> 126<212> DNA<213> Homo sapien
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60gattggctcg cagcaccac anaaaaggca gcagtggcag tggattgatg gggccatgta
120tttgta
126

<210> 1512<211> 331<212> DNA<213> Homo sapien
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331

<210> 1513<211> 350<212> DNA<213> Homo sapien
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350

<210> 1514<211> 170<212> DNA<213> Homo sapien
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170

<210> 1515<211> 174<212> DNA<213> Homo sapien
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174

<210> 1516<211> 481<212> DNA<213> Homo sapien
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480n
481

<210> 1517<211> 477<212> DNA<213> Homo sapien
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477

<210> 1518<211> 42<212> DNA<213> Homo sapien
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42

<210> 1519<211> 573<212> DNA<213> Homo sapien
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573

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571

<210> 1521<211> 117<212> DNA<213> Homo sapien
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117

<210> 1522<211> 123<212> DNA<213> Homo sapien
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123

<210> 1523<211> 461<212> DNA<213> Homo sapien
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461

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336

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234

<210> 1525<211> 438<212> DNA<213> Homo sapien
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180agagctgaga cgtacagtc cagtcttggga gatgacctg gactccatga gaaactcgaa
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438

<210> 1526<211> 308<212> DNA<213> Homo sapien
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180gtgtatgagg gggaaatggt ggggtcgtct gggccataga ggacattcag gatgactggg
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300cttgtgac
308

<210> 1527<211> 87<212> DNA<213> Homo sapien
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87

<210> 1528<211> 344<212> DNA<213> Homo sapien
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180gtgtatgagg gggaaatggt ggggtcgtct gggccataga ggacattcag gatgactggg
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300cttgtgacac tgagtagagt gaggtctgt ttgtcattgg acag
344

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180gtgaacctca gctctctgt coattgcagcc tctaacccac ctgcacagta ttcttgctg
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344

<210> 1530<211> 201<212> DNA<213> Homo sapien
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60gcccagagcc ctctccagag gttgggttga ccaactcact tggactcaga catatgaaga
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201

<210> 1531<211> 122<212> DNA<213> Homo sapien
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120ct
122

<210> 1532<211> 373<212> DNA<213> Homo sapien
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120tttccaggcg tccccagagg tctgtgcgac tagccctgt ctatcaaaag ttattagaga
180ggatgaagca ttactgtgaa gcactacagg aggaatgcac caggcagct ctcccacat
240ttctctcaga ttccacaga gactgtttga atgttttcaa aaccaagtat cacactttaa
300tgtacatggg ccgcaccata atgagatgtg agccttctgc atgtggggga ggagggagag

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235

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373

<210> 1533<211> 373<212> DNA<213> Homo sapien
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120aaatctgaga gaattggcgt gtagagtgcg gtggtgcatt cctctcctag tgtctcaagg
180taatgcttca tctctctcaa taacttttga tagacagggg ctagtgcgac agacotctgg
240gaagccctgg aaaacgctga tgcctgtttg aagatctcaa gcgcagagtc tgcaggttca
300tccctctctt cctgaggtct gttggtctga ggctgcagaa cattgtgtgat gacatggacc
360acgcatttg tgg
373

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120tttccagggc ttcccagagg tctgtgcgac tagccctgt ctatcaaaag ttattagaga
180ggatgaagca ttgacttgaa gcactacagg aggaatgcac caggcgagct ctccggcaat
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373

<210> 1535<211> 221<212> DNA<213> Homo sapien
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60caatccaaag cctggctcca gaagatcaca aagagccaaa gaaactggca ggtgtccatg
120cgctccaggg cagtgaatgt gttgtcaact actttttctg tgggaagaa ttatccatac
180ggaggatgct gaaggctcag agcttgaccc tgggccaact t
221

<210> 1536<211> 464<212> DNA<213> Homo sapien
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120ttaaaaataat gcttctcgat gtcagatttt acctgtttgc tgcgtagaaac atctctgctt
180aatttaccaa agccagacct tcaagtccaac atgcttctct agcttttcat agttgtctga
240catttccatg aaaacaaaag aaccaacttt gttttaacca aactttgttt gtttacagtt
300ttcaggggag cgtttctctc atgacacaca gcaacatccc aaagaaataa acaagtgtga
360caaaaaaaa acaaaccta aatgctactg ttccaaagag caacttgatg gtttttttta
420atactgagtg caaaaggtca cccaaattcc tatgatgaaa tttt 464

<210> 1537<211> 395<212> DNA<213> Homo sapien
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60ttctgtgtgga aaaaggcttt aatcctagaa tcttatatcc agccaaaatg gcaatttgatt
120ttagggggtaa aacaaaagta ttcttagta ttgaagaatt tagagattat gttttgcata
180tgccaccctt gagagaatta ctgggggaata atataccta gcaagccagg gtgaactaca
240acaactatgt ttctctcccc agcatgcata caaaaatcaa caagtataac gaaaataac
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360cagataatac atgatgtta ataaaggta tgttt 395

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120caatactctc acagcaacgt aggagctaga gctacatctt tcagaaacca aacttgctgt
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240gaacaagaag ttgaaggtta acatagaata ttattacag gcactcaac ctgcatttc
300ggaaggaat taggaatcca gatccgtga atttaactat tctgttacag cttgtcctgc
360aatctgctct ggagcaactt gctgcagag atttt 396

<210> 1539<211> 555<212> DNA<213> Homo sapien
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60ggagggaaga aggatactgt ggaagggat ggcggggcaa acatttagag ctagaagcca
120ctactggccc aatgctaaag ttctctgtct taagcctaaa aaagccagt tagtagggcc

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180cttatcactc ttagttttgct aggtttccccc tctgaaataa tgagcagatt tagccaggct
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 300ggtcggatct tgcaacggaa gtgttgccgc tlgttgcaat cgttgctgct ccaagtataa
 360aagttgttat tgagctcatc ctgcgcagct gctttgtccc acccatggac ttgcacagcc
 420aggtatctga cagatacatg gccocatcaa tccactgcc ctgctgctct tctgtgggt
 480gtgcagcgcc aatccatctc ggtcggcttc tctgatagcc acttatgtac tctgctatgg
 540tgcgtgcttc ctttt

555

<210> 1540<211> 358<212> DNA<213> Homo sapien
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 60ttaagactaa gtgtaactga caattgaatg aattaagcct aaaaacattt ctctaaagaa
 120ccagtggtctc atttaacatc ttgatgaac attattttaa tgactataaa aggatagtagc
 180agtatactga aattccactt aaatactgaa atattctact aatgacattt gttttgtcta
 240aatttctccc agaaaaatct gttagcattt cttaaaagtc cctcagattt gagggaaatt
 300ctaaattag acagttttct ctccaaataa atataaagta tcttgagtat ttttgttt

358

<210> 1541<211> 410<212> DNA<213> Homo sapien
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 120tgcagcgccg gaagtggggc agcagaggaga ggaagaggag catgcoccttc aaaaagggtg
 180cccgctttga gctggctctc atagtctcgg ctgagcacta caaggtgggtg gtaaatggaa
 240atccctctta tgagtacggg cacoggtctc cctacagat ggtcacccac ctgcagatgg
 300atgggggat gcaacttcaa tcaatcaact tcatcgagg ccagccocctc cggccccagg
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410

<210> 1542<211> 335<212> DNA<213> Homo sapien
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 120gcatgagtac catgtatttt ggaaaacttc cagtgttttt gtaagcttcc actgcggagg
 180gaaaaatgtaa aatgggggacc ccgaaataag tctgcatcat catcagtagc ctcaaatgtc
 240agacttccag gtgcactgag gggatggcag aagaacaagc ccgtgtatgc cttggctagc
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335

<210> 1543<211> 238<212> DNA<213> Homo sapien
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238

<210> 1544<211> 303<212> DNA<213> Homo sapien
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 180acactcagggt ggaggatttg ccccaaccag gactggcaaa ttaactttac tcaacatgac
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 303

300agg

<210> 1545<211> 276<212> DNA<213> Homo sapien
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276

<210> 1546<211> 344<212> DNA<213> Homo sapien
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 120ggaatatgcac ctccagacac agcaggagtc agcgggaggg cacagacatg cccctgcaca
 180ggcagaaaaat gggcctcctc aagcacaaaa gtgaccaagt acaatlttca gttgctaana
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344

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237

<210> 1547<211> 172<212> DNA<213> Homo sapien
 scocaaaaacag agtctctgggt gatatacatca tgaagccacg ctgtctctct ggatgggtttt
 60accacaagtc caattgtctat ggttaactca ggaactgtgag gaactggctct gatgcogagc
 120tcgagtgatca gtcttaacgga aacggagccc acctggcact tatcttgagt tt 172

<210> 1548<211> 1071<212> DNA<213> Homo sapien
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 900aattctgtttt gtttttagtt gtctctcaaga gacaacagct tcaagtaatt tctcatgat
 960ttgggtgtgg ctaagctggg gattggttct gttccctctg cctcgtgtga gagaaaagct
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<210> 1549<211> 539<212> DNA<213> Homo sapien
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 120tctcaactct tgtgtgaga aggttatga ctgacctga atcagtttcc
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<210> 1550<211> 520<212> DNA<213> Homo sapien
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 480gtgggttggga ttgactggcc taccttggtc actctttaat 520

<210> 1551<211> 340<212> DNA<213> Homo sapien
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 180gaaggcagga gccccaagctg gaacctcogag ggccatgag aggcacgacg actcttggcc
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 300cagactcag gaagagaaga gctggggagc aggtgaaag 340

<210> 1552<211> 1072<212> DNA<213> Homo sapien
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360gtgtcattga agatttcagt taotacacta ggcactgaag taccattctg gagggctgtc
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 840cttgctaaagc aaagttagoca tgcctctccg tctatggtct attctgttta tctttccacc
 900aatgaatact ctgatcttacc agtctttcca tttttctctg gtctgcagaa ggtaagggtat
 960caataaggct aaacctccat catcaaaaag ccaccagaca tcaatagtat tcttctctgt
 1020tttttctgta aactgtgtac tagcttcaag aagcttttgg tcatgtacat tt

1072

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 120gtcaccacaa agctcagagc ctggaagccc agatagacaa cagagatgct tccagaattt
 180aacccccaact ccatcctatg ctgctacaat tgcttctaaa ctggcaattc .acaatccaaa
 240ttttaccacc accctgccag ttaactcaca aaacatccaa cctgtcagat acaatagaa
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384

<210> 1554<211> 408<212> DNA<213> Homo sapien
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 300gtttttctgt ccttttgctc ccgggaagcg cttctgtctga agttctatc ctggagcctg
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408

<210> 1555<211> 607<212> DNA<213> Homo sapien
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 180ggaaatgtcca gggaacctc ttcaaagtga ttactaagga tgacacacac tattacattc
 240aggccagcag caaggtctgag cgagccagat ggattgaagc tatcaaaaag ctacatgac
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 420aatgtctgcca agagtctctc agattacaaa cagcagtggt gccatttctc tcccatctt
 480catgttacaa acctggaaa gctagaacag ccattaggcg tcagcatctt gacttttccc
 540cgcatacaca aacagccatt tctcgggca ccaaatgtag ttccctttgt tgaacaatt
 600acactgg

607

<210> 1556<211> 192<212> DNA<213> Homo sapien
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 60aaaagaggaa tttccagatg gcagctactg gctcagaacc agggggtccc ttgccaagtc
 120gtctctctatg tggctcccg aattgtctgag gtctcaact tcagagggtc ttgatggaaa
 180ataaagcaac ag

192